

181150

From: Whiteman, Brian  
Sent: Friday, March 03, 2006 9:19 AM  
To: STIC-Biotech/ChemLib  
Subject: seq search

10/782,899

nucleotides 1-192 of SEQ ID NO: 1 and nucleotides 472-600 of SEQ ID NO: 1  
1) us patents and published us patent application databases

Thank you,

Brian Whiteman  
Remsen, 2D14  
mail box 2C18  
Patent Examiner - Art Unit 1635  
United States Patent and Trademark Office  
(571) 272-0764

RECEIVED  
MAR - 3 2006  
STIC

\*\*\*\*\*

Searcher: \_\_\_\_\_  
Searcher Phone: \_\_\_\_\_  
Date Searcher Picked up: \_\_\_\_\_  
Date completed: \_\_\_\_\_  
Searcher Prep Time: \_\_\_\_\_  
Online Time: \_\_\_\_\_

\*\*\*\*\*

Type of Search  
NA# \_\_\_\_\_ AA# \_\_\_\_\_  
S/L: \_\_\_\_\_ Oligomer: \_\_\_\_\_  
Encode/Transl: \_\_\_\_\_  
Structure #: \_\_\_\_\_ Text: \_\_\_\_\_  
Inventor: \_\_\_\_\_ Litigation: \_\_\_\_\_

\*\*\*\*\*

Vendors and cost where applicable  
STN: \_\_\_\_\_  
DIALOG: \_\_\_\_\_  
QUESTEL/ORBIT: \_\_\_\_\_  
LEXIS/NEXIS: \_\_\_\_\_  
SEQUENCE SYSTEM: \_\_\_\_\_  
WWW/Internet: \_\_\_\_\_  
Other (Specify): \_\_\_\_\_

**This Page Blank (uspto)**

GenCore version 5.1.7  
Copyright (c) 1993 - 2006 Bioceleration Ltd.

OM nucleic - nucleic search, using sw model

Run on: March 7, 2006, 11:25:13 ; Search time 1781.23 Seconds  
(without alignments)  
6127.182 Million cell updates/sec

Title: US-10-782-899-1\_COPY\_1\_192

Perfect score: 192  
Sequence: 1 gctggcgccggcccatga.....tgaccacaagaagatgcttt 192

Scoring table: IDENTITY NUC  
Gapop 10.0 , Gapext 1.0

Searched: 5883141 seqs, 28421725653 residues

Total number of hits satisfying chosen parameters: 1176282

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

- 1: gb\_ba.\*
- 2: gb\_in.\*
- 3: gb\_env.\*
- 4: gb\_ov.\*
- 5: gb\_ov.\*
- 6: gb\_pat.\*
- 7: gb\_ph.\*
- 8: gb\_pr.\*
- 9: gb\_ro.\*
- 10: gb\_sts.\*
- 11: gb\_sy.\*
- 12: gb\_un.\*
- 13: gb\_vi.\*
- 14: gb\_htg.\*
- 15: gb\_pl.\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	192	100.0	600	6	BD144660 Drug for
2	175.2	91.2	56640	1	Continuation (23 o
3	175.2	91.2	156638	6	AX492788 Sequence
4	175.2	91.2	156646	6	AX553955 Sequence
5	173.2	90.2	600	1	AB072446 Bifidobac
6	37	19.3	230625	14	AC096816 Rattus no
7	37	19.3	246991	14	AC122956 Rattus no
8	36.4	19.0	67395	8	AL671518 Human DNA
9	36.4	19.0	317403	14	AC154842 Mus muscu
10	35.8	18.6	170350	8	AC140952
11	35.8	18.6	171239	14	AC092873 Pan trogl
12	35.8	18.6	171391	14	AC098646 Pan trogl
13	35.8	18.6	183888	8	AC151799 Pan trogl
14	35	18.2	181622	14	AC149460 Papio anu
15	35	18.2	231844	14	AC116068 Rattus no
16	35	18.2	234234	14	AC097194 Rattus no
17	34.8	18.1	152430	8	AC087432 Homo sapi
18	34.8	18.1	160410	14	AC020753 Homo sapi

19	34.8	18.1	166162	8	AC018814
20	34.8	18.1	181870	14	AC024216
21	34.8	18.1	188432	14	AC018497
22	34.4	17.9	148269	14	AC026486
23	34.4	17.9	162249	14	AC020682
24	34.4	17.9	166973	8	AL451049
25	34.2	17.8	155711	2	VYIVD10
26	34.2	17.8	183089	9	AL663115
27	34.2	17.8	189862	14	AC157771
28	34.2	17.8	213720	9	AC141477
29	34.2	17.8	235639	9	AC158618
30	34.2	17.7	166873	8	AC008554
31	34	17.7	179393	8	AC010636
32	33.6	17.5	4799	9	AF367244
33	33.6	17.5	17073	2	AC084160
34	33.6	17.5	179650	9	AC158525
35	33.4	17.4	114784	8	HS118924
36	33.2	17.3	630	10	G54937
37	33.2	17.3	4765	9	BC052387
38	33.2	17.3	5011	9	AF140786
39	33.2	17.3	5034	9	AF387811
40	33.2	17.3	74436	14	AC026625
41	33.2	17.3	87258	8	AF091512
42	33.2	17.3	110000	15	CR382122_05
43	33.2	17.3	132910	14	AC014319
44	33.2	17.3	160421	8	AC116546
45	33.2	17.3	168700	14	AC009115

ALIGNMENTS

RESULT 1  
BD144660  
LOCUS Drug for gene therapy using anaerobic bacteria.  
DEFINITION Drug for gene therapy using anaerobic bacteria.  
ACCESSION BD144660  
VERSION BD144660.1 GI:27850418  
KEYWORDS JP 2002097144-A/1.  
SOURCE Bifidobacterium longum  
ORGANISM Bifidobacterium longum  
Bacteria; Actinobacteria; Actinobacteridae; Bifidobacteriales;  
Bifidobacteriaceae; Bifidobacterium.  
REFERENCE 1 (bases 1 to 600)  
AUTHORS Fujimori,M., Taniguchi,S., Amano,J., Yazawa,K., Kano,Y.,  
Nakamura,T. and Sasaki,T.  
TITLE Drug for gene therapy using anaerobic bacteria  
JOURNAL Patent: JP 2002097144-A 1 02-APR-2002;  
JUN AMANO,MINORU FUJIMORI  
COMMENT OS Bifidobacterium longum  
PN JP 2002097144-A/1  
PD 02-APR-2002  
PF 21-SEP-2001 JP 2001290187  
PI MINORU FUJIMORI,SHUNICHIRO TANIGUCHI,JUN  
AMANO,KAZUTORA YAZAWA,  
PI YASUMASA KANO,TOSHIIYUKI NAKAMURA,TAKAYUKI SASAKI PC  
A61K35/74,A61K31/396,A61K31/7068,A61K31/711,A61K38/00,A61K38/00 PC  
43,A61K48/00,  
PC A61P35/00,C12N1/21,C12N15/09//C12N1/21,C12N1/01,C12N15/00,  
PC A61K37/02,  
PC A61K37/48  
CC Drug for gene therapy using anaerobic bacteria FH Key  
Location/Qualifiers  
FT CDS Location/Qualifiers  
(193)..(471).

FEATURES  
source  
1..600  
/organism="Bifidobacterium longum"  
/mol\_type="genomic DNA"  
/db\_xref="taxon:216816"

ORIGIN

Query Match 100.0%; Score 192; DB 6; Length 600;  
Best Local Similarity 100.0%; Pred. No. 1.4e-49;

Matches	192;	Conservative	0;	Mismatches	0;	Indels	0;	Gaps	0;
Qy	1	GCTGGCGCGGGCCATGAAGTGGCTTGACAAAGCATAACTCTGTCTGATTGCTCTATTT	60						
Db	1	GCTGGCGCGGGCCATGAAGTGGCTTGACAAAGCATAACTCTGTCTGATTGCTCTATTT	60						
Qy	61	TCATACTCTCGGGGAATAGATGTGMAAACCTTTATAAAGCGGGGTTTTTCGCAGAAAC	120						
Db	61	TCATACTCTCGGGGAATAGATGTGMAAACCTTTATAAAGCGGGGTTTTTCGCAGAAAC	120						
Qy	121	ATGCGCTAGTATCATTTGATGACCAACATGGACATGAACAAAGTGCTTGTCCTCTGACCCAA	180						
Db	121	ATGCGCTAGTATCATTTGATGACCAACATGGACATGAACAAAGTGCTTGTCCTCTGACCCAA	180						
Qy	181	GAAGGATGCTTT	192						
Db	181	GAAGGATGCTTT	192						

RESULT 2  
AE014295\_22/c  
WPCOMMENT

Sequence split into 23 fragments LOCUS AE014295 Accession AE014295

Fragment Name	Begin	End
AE014295_00	1	110000
AE014295_01	100001	210000
AE014295_02	200001	310000
AE014295_03	300001	410000
AE014295_04	400001	510000
AE014295_05	500001	610000
AE014295_06	600001	710000
AE014295_07	700001	810000
AE014295_08	800001	910000
AE014295_09	900001	1010000
AE014295_10	1000001	1110000
AE014295_11	1100001	1210000
AE014295_12	1200001	1310000
AE014295_13	1300001	1410000
AE014295_14	1400001	1510000
AE014295_15	1500001	1610000
AE014295_16	1600001	1710000
AE014295_17	1700001	1810000
AE014295_18	1800001	1910000
AE014295_19	1900001	2010000
AE014295_20	2000001	2110000
AE014295_21	2100001	2210000
AE014295_22	2200001	2256640

Query Match	91.2%	Score 175.2;	DB 1;	Length 56640;
Best Local Similarity	97.4%;	Pred. No. 2.6e-44;		
Matches 189;	Conservative 0;	Mismatches 3;	Indels 2;	Gaps 1;
QY	1	GC TGGCGCGCGCGCCATGAAGTGGCTTGAACAAGCATAAATCTTGCTGTGATTCGCTCTATT	60	
Db	41859	GCTGGCGCGCGCGCCATGAGTGGCTTGACAAGCATAAATCTTGCTGTGATTCGCTCTATT	41800	
QY	61	TC--ANATCCTTCGGGGAAATAGATGTGAAAAACCTTATAAAACGCGGGTTTTTCGAGAA	118	
Db	41799	TCAATACCCCTTCGGGGAAATAGATGTGAAAAACCTTATAAAACGCGGGTTTTTCGAGAA	41740	
QY	119	ACATGGCGCTAGTATCATTTGATGACAACATGGACTAAGCAAAAGTGCTGCTGCCCTTGACCC	178	
Db	41739	ACATGGCGCTAGTATCATTTGATGACAACATGGACTAAGCAAAAGTGCTGCTGCCCTTGACCC	41680	
QY	179	AGAAGGATGCTTT	192	
Db	41679	AGAAGGATGCTTT	41666	

RESULT 3	DNA	linear	PAT 26-SEP-2002
AX492788/c	156638 bp		
LOCUS	AX492788		

DEFINITION	Sequence 1106 from Patent EP12227152.
ACCESSION	AX492788
VERSION	AX492788.1 GI:23338471
KEYWORDS	
SOURCE	Bifidobacterium longum biovar Longum
ORGANISM	Bifidobacterium longum biovar Longum
	Bacteria; Actinobacteria; Actinobacteriales; Bifidobacteriales; Bifidobacteriaceae; Bifidobacterium.
REFERENCE	1
AUTHORS	None.
TITLE	Bacterial strain and genome of Bifidobacterium
JOURNAL	Patent: EP 1227152-A 1106 31-JUL-2002;
	Societe des Produits Nestle S.A. (CH)
FEATURES	Location/Qualifiers
source	1..156638
	/organism="Bifidobacterium longum biovar Longum"
	/mol_type="unassigned DNA"
	/db_xref="taxon:1673"
	/note="1106 seq in place of 1099 because seq 1 (2.256.638
	bases splitted
	in 7 more sequences.
	seq 0001: from 0.000.001 to 0.349.980 length: 349980
	seq 1100: from 0.300.001 to 0.649.980 length: 349980
	seq 1101: from 0.600.001 to 0.949.980 length: 349980
	seq 1102: from 0.900.001 to 1.249.980 length: 349980
	seq 1103: from 1.200.001 to 1.549.980 length: 349980
	seq 1104: from 1.500.001 to 1.849.980 length: 349980
	seq 1105: from 1.800.001 to 2.149.980 length: 349980
	seq 1106: from 2.100.001 to 2.256.638 length: 156638"
ORIGIN	
Query Match	91.2%; Score 175.2; DB 6; Length 156638;
Best Local Similarity	97.4%; Pred. No. 2.6e-44;
Matches 189; Conservative	0; Mismatches 3; Indels 2; Gaps 1;
Qy	1 GCTGGCGCGCGCCATGAAGTGGCTTGACAGCATATCTTGTCTGATTCGTCTATT 60
Db	142164 GCTGGCGCGCGCCATGAAGTGGCTTGACAGCATATCTTGTCTGATTCGTCTATT 142105
Qy	61 TC--AATACCTTCGGGGAATAGATGTGMAAACCTTTATAAAACCGGGTTTCGCAGAA 118
Db	142104 TCATACCCCTTCGGGGAATAGATGTGMAAACCTTTATAAAACCGGGTTTCGCAGAA 142045
Qy	119 ACATGCGCTAGTATCATTTGATGACACATGGACTAAGCAAAAGTCTTGTCCCTGACCC 178
Db	142044 ACATGCGCTAGTATCATTTGATGACACATGGACTAAGCAAAAGTCTTGTCCCTGACCC 141985
Qy	179 AAGAAGGATGCTTT 192
Db	141984 AAGAAGGATGCTTT 141971
RESULT 4	
AX553955/c	
LOCUS	AX553955 156646 bp DNA linear PAT 27-NOV-2002
DEFINITION	Sequence 9 from Patent WO02074798.
ACCESSION	AX553955
VERSION	AX553955.1 GI:25897908
KEYWORDS	
SOURCE	Bifidobacterium longum biovar Longum
ORGANISM	Bifidobacterium longum biovar Longum
	Bacteria; Actinobacteria; Actinobacteriales; Bifidobacteriales; Bifidobacteriaceae; Bifidobacterium.
REFERENCE	1
AUTHORS	Arigoni, F., Delley, M., Mollet, B., Pridmore, R.D., Schell, M.A., Pohl, T.G. and Zwhalen, M.C.
TITLE	The genome of a bifidobacterium
JOURNAL	Patent: WO 02074798-A 9 26-SEP-2002;
	SOCIETE DES PRODUITS NESTLE S.A. (CH)
FEATURES	Location/Qualifiers
source	1..156646
	/organism="Bifidobacterium longum biovar Longum"
	/mol_type="unassigned DNA"



/db\_xref="taxon:1679"  
/note="seq 1: original length 2256646 splitted into  
seq 1: from 0.000.001 to 0.349.980  
seq 3: from 0.300.001 to 0.649.980  
seq 4: from 0.600.001 to 0.949.980  
seq 5: from 0.900.001 to 1.249.980  
seq 6: from 1.200.001 to 1.549.980  
seq 7: from 1.500.001 to 1.849.980  
seq 8: from 1.800.001 to 2.149.980  
seq 9: from 2.100.001 to 2.256.646"

ORIGIN

Query Match 91.2%; Score 175.2; DB 6; Length 156646;  
Best Local Similarity 97.4%; Pred. No. 2.6e-44;  
Matches 189; Conservative 0; Mismatches 3; Indels 2; Gaps 1;

QY 1 GCTGGCGCGCGCCATGAAGTGGCTTGACAAGCATAATCTTGTCTGATTCGCTCTATT 60  
DB 142172 GCTGGCGCGCGCCATGAAGTGGCTTGACAAGCATAATCTTGTCTGATTCGCTCTATT 142113

QY 61 TC--AATACCTTCGGGGAAATAGATGTGAAAACCCCTTATAAAACGGCGGTTTCGCAGAA 118  
DB 142112 TCATACCCCTTCGGGGAAATAGATGTGAAAACCCCTTATAAAACGGCGGTTTCGCAGAA 142053

QY 119 ACATGGCTAGTATCATGTGATGACACATGAGCAAGCAAAAGTGTGTCCTGACCC 178  
DB 142052 ACATGGCTAGTATCATGTGATGACACATGAGCAAGCAAAAGTGTGTCCTGACCC 141993

QY 179 AAGAAGGATGCTTT 192  
DB 141992 AAGAAGGATGCTTT 141979

RESULT 5  
AB072446  
LOCUS  
DEFINITION  
ACCESSION  
VERSION  
KEYWORDS  
SOURCE  
ORGANISM  
REFERENCE  
AUTHORS  
TITLE  
JOURNAL  
PUBMED  
REFERENCES  
AUTHORS  
TITLE  
JOURNAL

AB072446  
Bifidobacterium longum biovar Longum hup gene for histone-like HU  
protein HbL, complete cds.  
AB072446  
AB072446.1 GI:19918953  
Bifidobacterium longum biovar Longum  
Bifidobacterium longum biovar Longum  
Bacteria; Actinobacteria; Actinobacteridae; Bifidobacteriales;  
Bifidobacteriaceae; Bifidobacterium.  
1  
Takeuchi, A., Matsumura, H. and Kano, Y.  
Cloning and expression in Escherichia coli of a gene, hup, encoding  
the histone-like protein HU of Bifidobacterium longum  
Biosci. Biotechnol. Biochem. 66 (3), 598-603 (2002)  
12005055  
2 (bases 1 to 600)  
Kano, Y.  
Direct Submission  
Submitted (03-Oct-2001) Yasunobu Kano, Kyoto Pharmaceutical  
University, Institute of Molecular and Cellular Biology for  
Pharmaceutical Sciences; 1, Shichono-cho, Misasagi, Yamashina-ku,  
Kyoto 6078412, Japan (E-mail: ykano@mb.kyoto-phu.ac.jp,  
Tel: 81755954719, Fax: 81755954799)

FEATURES  
Source  
RBS  
gene  
CDS

1. .600  
/organism="Bifidobacterium longum biovar Longum"  
/mol\_type="genomic DNA"  
/db\_xref="taxon:1679"  
180\_ .185  
193\_ .474  
/genes="hup"  
/genes="hup"  
/codon\_start=1  
/translation=11  
/product="histone-like HU protein HbL"  
/protein\_id="BAB88690.1"

/db\_xref="GI:19918954"  
/translation="WAYNKSQDLVSKIAQKSNLTKAQAEAAVNAFQDFVFAAMKSGEGL  
KLTLGFSAEVRKPRTPGRNPRTEGQIDIPASYGVRSISAGSLKKAATVE"

ORIGIN

Query Match 90.2%; Score 173.2; DB 1; Length 600;  
Best Local Similarity 97.4%; Pred. No. 1.3e-43;  
Matches 187; Conservative 0; Mismatches 3; Indels 2; Gaps 1;

QY 3 TGGGCGCGCGCCATGAAGTGGCTTGACAAGCATAATCTTGTCTGATTCGCTCTATTTC 62  
DB 1 TGGGCGCGCGCCATGAAGTGGCTTGACAAGCATAATCTTGTCTGATTCGCTCTATTTC 60

QY 63 --AATACCTTCGGGGAAATAGATGTGAAAACCCCTTATAAAACGGCGGTTTCGAGAAAC 120  
DB 61 ATACCCCTTCGGGGAAATAGATGTGAAAACCCCTTATAAAACGGCGGTTTCGAGAAAC 120

QY 121 ATGCGCTAGTATCATGTGATGACACATGAGCAAGCAAAAGTGTGTCCTGACCCGACCCAA 180  
DB 121 ATGCGCTAGTATCATGTGATGACACATGAGCAAGCAAAAGTGTGTCCTGACCCGACCCAA 180

QY 181 GAAGGATGCTTT 192  
DB 181 GAAGGATGCTTT 192

RESULT 6  
AC096816/c  
LOCUS  
DEFINITION  
ACCESSION  
VERSION  
KEYWORDS  
SOURCE  
ORGANISM  
REFERENCE  
AUTHORS

AC096816  
Rattus norvegicus clone CH230-171E19, \*\*\* SEQUENCING IN PROGRESS  
\*\*\*, 7 unordered pieces.  
AC096816  
AC096816.6 GI:30520628  
HTG; HTGS PHASE1; HTGS DRAFT; HTGS\_ENRICHED.  
Rattus norvegicus (Norway rat)  
Rattus norvegicus  
Rattus norvegicus  
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
Mammalia; Eutheria; Euarchontoglires; Glires; Rodentia;  
Sciurognathi; Muridea; Muridae; Murinae; Rattus.  
1 (bases 1 to 230625)  
Muzny, D., Maric, Metzker, M., Lee, A., Adams, C., Alder, J.,  
Allen, C., Allen, H., Altschuler, S., Amin, A., Anguiano, D.,  
Anyalebech, V., Aoyagi, A., Ayodeji, M., Baca, E., Baden, H.,  
Baldwin, D., Bandaranaike, D., Barber, M., Barnstead, M., Benahmed, F.,  
Biswal, K., Blair, J., Blankenburg, K., Blyth, P., Brown, M.,  
Bryant, N., Buhay, C., Burch, P., Burrell, K., Calderon, E.,  
Cardenas, V., Carter, K., Cavazos, I., Caesar, H., Center, A.,  
Chacko, J., Chavez, D., Chen, G., Chen, R., Chen, Y., Chen, Z., Chu, J.,  
Cleveland, C., Cockrell, R., Cox, C., Coyle, M., Cree, A., D'Souza, L.,  
Davila, M., Davis, C., Davy-Carroll, L., De Anda, C., Dederich, D.,  
Delgado, O., Denson, S., Deramo, C., Ding, Y., Dinh, H., Divya, K.,  
Draper, H., Dugan-Rocha, S., Dunn, A., Durbin, K., Duval, B., Eaves, K.,  
Egan, A., Escoto, M., Eugene, C., Evans, C., Falls, T., Fan, G.,  
Fernandez, S., Finley, M., Flagg, N., Forbes, L., Foster, M., Foster, P.,  
Fraser, C., Gabis, A., Ganta, R., Garcia, A., Garner, T., Garza, M.,  
Gebrgeorgis, E., Geer, K., Gill, R., Grady, M., Guerra, W., Guervara, W.,  
Gunaratne, P., Haaland, W., Hamill, C., Hamilton, C., Hamilton, K.,  
Harvey, Y., Havlak, P., Hawes, A., Henderson, N., Hernandez, J.,  
Hernandez, R., Hines, S., Hladun, S., Hodgson, A., Hughes, M.,  
Hollins, B., Howells, S., Hulyk, S., Hume, J., Idelbird, D., Jackson, A.,  
Jackson, L., Jacob, L., Jiang, H., Johnson, B., Johnson, R., Jolivet, A.,  
Karpach, S., Kelly, S., Kelly, S., Khan, Z., King, L., Kovar, C.,  
Kowals, C., Kraft, C., Lebow, H., Levan, J., Lewis, L., Li, Z., Liu, J.,  
Liu, J., Liu, W., Liu, Y., London, P., Longacre, S., Lopez, J.,  
Lorensuhewa, L., Loulsegged, H., Lozada, R., Lu, X., Ma, J.,  
Maheshwari, M., Mahindartne, M., Mahmoud, M., Malloy, K., Mangum, A.,  
Mangum, B., Mapua, P., Martin, K., Martin, R., Martinez, E.,  
Mawhiney, S., McLeod, M., McNeill, T., Meenan, E.,  
Milosavljevic, A., Miner, G., Minja, E., Montemayor, J., Moore, S.,  
Morgan, M., Morris, K., Morris, S., Munidasa, M., Murphy, M., Nair, L.,  
Nankervis, C., Neal, D., Newton, N., Nguyen, N., Norrie, S., Parks, K.,  
Paokelemeh, O., Okwuonu, G., Olarnpunsagoon, A., Pal, S., Parks, K.,  
Pasternak, S., Paul, H., Perez, A., Perez, L., Pfannkuch, C.,

Plopper, F., Poindexter, A., Popovic, D., Primus, E., Pu, L.-L.,  
 Puazo, M., Quiroz, J., Rachlin, E., Reeves, K., Regier, M.A., Reigh, R.,  
 Reilly, B., Reilly, M., Ren, Y., Reuter, M., Richards, S., Riggs, F.,  
 Rives, C., Rodkey, T., Rojas, A., Rose, M., Rose, R., Ruiz, S.J.,  
 Sanders, W., Savery, G., Scherer, S., Scott, G., Shatsman, S., Shen, H.,  
 Shetty, J., Shvartbeyn, A., Sleson, I., Sitter, C.D., Smaje, D.,  
 Sneed, A., Sodergren, E., Song, X.-Z., Sorelle, R., Sosa, J.,  
 Steinle, M., Strong, R., Sutton, A., Svatek, A., Tabor, P., Taylor, C.,  
 Taylor, T., Thomas, N., Thomas, S., Tingey, A., Trejos, Z., Usmani, K.,  
 Valas, R., Vera, V., Villalana, D., Waldron, L., Walker, B., Wang, J.,  
 Wang, Q., Wang, S., Warren, J., Warren, R., Wei, X., White, F.,  
 Williams, G., Willson, R., Wleczek, R., Wooden, H., Worley, K.,  
 Wright, D., Wright, R., Wu, J., Yakub, S., Yen, J., Yoon, L., Yoon, V.,  
 Yu, F., Zhang, J., Zhou, J., Zhou, X., Zhao, S., Dunn, D., von  
 Niederhausern, A., Weiss, R., Smith, D.R., Holt, R.A., Smith, H.O.,  
 Weinstock, G. and Gibbs, R.A.

# TITLE

Unpublished

2 (bases 1 to 230625)

Worley, K.C.

Direct Submission

Submitted (28-SEP-2001) Human Genome Sequencing Center, Department  
 of Molecular and Human Genetics, Baylor College of Medicine, One  
 Baylor Plaza, Houston, TX 77030, USA

3 (bases 1 to 230625)

Rat Genome Sequencing Consortium.

Direct Submission

Submitted (10-MAY-2003) Human Genome Sequencing Center, Department  
 of Molecular and Human Genetics, Baylor College of Medicine, One  
 Baylor Plaza, Houston, TX 77030, USA

## COMMENT

On May 10, 2003 this sequence version replaced gi:24942762.  
 The sequence in this assembly is a combination of BAC based reads  
 and whole genome shotgun sequencing reads assembled using Atlas  
 (<http://www.hgsc.bcm.tmc.edu/projects/rat/>). Each contig described  
 in the feature table below represents a scaffold in the Atlas  
 assembly (a 'contig-scaffold'). Within each contig-scaffold,  
 individual sequence contigs are ordered and oriented, and separated  
 by sized gaps filled with Ns to the estimated size. The sequence  
 may extend beyond the ends of the clone and there may be sequence  
 contigs within a contig-scaffold that consist entirely of whole  
 genome shotgun sequence reads. Both end sequences and whole genome  
 shotgun sequence only contigs will be indicated in the feature  
 table.

----- Genome Center

Center: Baylor College of Medicine

Center code: BCM

Web site: <http://www.hgsc.bcm.tmc.edu/>

Contact: hgsc-help@bcm.tmc.edu

----- Project Information

Center project name: GFJV

Center clone name: CH230-171E18

----- Summary Statistics

Assembly program: Atlas 3.0;

Consensus quality: 197146 bases at least Q40

Consensus quality: 202146 bases at least Q30

Consensus quality: 205589 bases at least Q20

Estimated insert size: 207699; sum-of-contigs estimation

Quality coverage: 6x in Q20 bases; sum-of-contigs estimation

-----

\* NOTE: Estimated insert size may differ from sequence length  
 (see [http://www.hgsc.bcm.tmc.edu/docs/genbank\\_draft\\_data.html](http://www.hgsc.bcm.tmc.edu/docs/genbank_draft_data.html)).

\* NOTE: This is a 'working draft' sequence. It currently

\* consists of 7 contigs. The true order of the pieces

\* is not known and their order in this sequence record is

\* arbitrary. Gaps between the contigs are represented as

\* runs of N, but the exact sizes of the gaps are unknown.

\* This record will be updated with the finished sequence

\* as soon as it is available and the accession number will

\* be preserved.

\* 1 6876: contig of 6876 bp in length

\* 6877 6976: gap of unknown length

\* 6977 35044: contig of 28068 bp in length

\* 35045 35144: gap of unknown length

\* 35145 60636: contig of 25492 bp in length

\* 60637 60736: gap of unknown length

\* 60737 72181: contig of 11445 bp in length

\* 72182 72281: gap of unknown length

\* 72282 227314: contig of 15033 bp in length

\* 227315 227414: gap of unknown length

\* 227415 228683: contig of 1269 bp in length

\* 228684 228783: gap of unknown length

\* 228784 230625: contig of 1842 bp in length.

## FEATURES

source

1. .230625

/organism="Rattus norvegicus"

/mol\_type="genomic DNA"

/db\_xref="taxon:10116"

/clone="CH230-171E18"

6877..6976

/estimated\_length=unknown

35045..35144

/estimated\_length=unknown

60637..60736

/estimated\_length=unknown

72182..72281

/estimated\_length=unknown

224079..227314

/notes="wgs contig"

227315..227414

/estimated\_length=unknown

228684..228783

/estimated\_length=unknown

## ORIGIN

Query Match 19.1%; Score 37; DB 14; Length 230625;

Best Local Similarity 54.9%; Pred.No. 1.5;

Matches 73; Conservative 0; Mismatches 60; Indels 0; Gaps 0;

QY 29 GACAAGCATATCTTGTCTGATTCGTCATATTTTCAATACCTTCGGGAAATAGATGTGAA 88

Db 127530 GAAAACCCATTATGTCTAAGAGCTCTATGCAGATATCTAATCTCTAAAGGAACATAAA 127471

QY 89 AACCTTATAAAGCGGGTTTCGCGAAGAACATCGCTAGTATCATTCATGATGACAAATG 148

Db 127470 AATACTTTTCAACATGATATTTTCTTAGTAGGAGTAGTATCTTCTTAACCCCTAG 127411

QY 149 GACTAGCAAAAG 161

Db 127410 GACTAGCAGAGG 127398

## RESULT 7

AC122956/c

LOCUS

AC122956

DEFINITION

Rattus norvegicus clone CH230-17E8, WORKING DRAFT SEQUENCE.

AC122956

ACCESSION

AC122956.4

VERSION

HTG; HTGS PHASE2; HTGS DRAFT; HTGS\_FULLTOP.

KEYWORDS

Rattus norvegicus (Norway rat)

SOURCE

ORGANISM

Rattus norvegicus

Sukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

Mammalia; Eutheria; Euarchontoglires; Glires; Rodentia;

Sciurognathi; Muridae; Murinae; Rattus.

REFERENCE

1 (bases 1 to 246991)

AUTHORS

Muzny, D., Marle, Metzker, M., Lee, S., Abramson, S., Adams, C., Alder, J.,

Allen, C., Allen, H., Alebrooks, S., Amin, A., Angiano, D.,

Anyaiebechi, V., Aoyagi, A., Ayodeji, M., Baca, E., Baden, H.,

Baldwin, D., Bandaranaike, D., Barber, M., Barnstead, M., Benahmed, F.,

Blawie, K., Blair, J., Blankenburg, K., Blyth, P., Brown, M.,

Bryant, N., Buhay, C., Burch, P., Burrell, K., Calderon, E.,

Cardenas, V., Carter, K., Cavazos, I., Cesar, H., Chen, A.,

Chacko, J., Chavez, D., Chen, G., Chen, Y., Chen, Z., Chu, J.,

Cleveland, C., Cockrell, R., Cox, C., Coyle, M., Cree, A., D'Souza, L.,

Davila, M.L., Davis, C., Davy-Carroll, L., De Anda, C., Dederich, D.,

Delgado, O., Denson, S., Deramo, C., Ding, Y., Dinh, H., Divya, K.,

Draper, H., Dugan-Rocha, S., Dunn, A., Durbin, K., Duval, B., Eaves, K.,

Egan, A., Escotto, M., Eugene, C., Evans, C.A., Falls, T., Fan, G., Fernandez, S., Finley, M., Flagg, N., Forbes, L., Foster, M., Foster, P., Fraser, C.M., Gaisi, A., Ganta, R., Garcia, A., Garner, T., Garza, M., George, P., Geer, K., Gill, R., Grady, M., Guerra, W., Guevara, W., Gunaratne, P., Haaland, W., Hamil, C., Hamilton, C., Hamilton, K., Harvey, Y., Havlak, P., Haves, A., Henderson, N., Hernandez, J., Hernandez, R., Hines, S., Hladun, S.L., Hodgson, A., Hogues, M., Hollins, B., Howells, S., Hulyk, S., Hume, J., Idlebird, D., Jackson, A., Jackson, L., Jacob, L., Jiang, H., Johnson, B., Johnson, R., Jolivet, A., Karpach, S., Kelly, S., Kelly, S., Khan, Z., King, L., Kovar, C., Kowis, C., Kraft, C.L., Lebow, H., Levan, J., Lewis, L., Li, Z., Liu, J., Liu, J., Liu, W., Liu, Y., London, P., Longacre, S., Lopez, J., Lorenshewa, L., Louised, H., Lozano, R.J., Lu, X., Ma, J., Maheshwari, M., Mahindartne, M., Mahmoud, M., Malloy, K., Mangum, A., Mangum, B., Mapua, P., Martin, K., Martin, R., Martinez, E., Mawhney, S., McLeod, M.P., McNeill, T.Z., Meenen, E., Milosavljevic, A., Miner, G., Minja, E., Montemayor, J., Moore, S., Morgan, M., Morris, K., Morris, S., Munidasa, M., Murphy, M., Nair, L., Nwaokeme, O., Okwuonu, G., Olarnpunsagoon, A., Pal, S., Parks, K., Pasternak, S., Paul, H., Perez, A., Perez, L., Pfannkuch, C., Plopper, F., Poindexter, A., Popovic, D., Primus, E., Pu, L., Puazo, M., Quiroz, J., Rachlin, E., Reeves, K., Regier, M.A., Reich, R., Reilly, B., Reilly, M., Ren, Y., Reuter, M., Richards, S., Riggs, P., Rives, C., Rodkey, T., Rojbas, A., Rose, M., Rose, R., Ruiz, S.J., Sanders, W., Savary, G., Scherer, S., Scott, G., Shatsman, S., Shen, H., Shetty, J., Shvartsbeyn, A., Sisson, I., Sitter, C.D., Smajs, D., Sneed, A., Sodergren, E., Song, X.-Z., Sorelle, R., Soosa, J., Steidle, M., Strong, R., Sutton, A., Svatek, A., Tabor, P., Taylor, C., Taylor, T., Thomas, N., Thomas, S., Tingey, A., Trejos, Z., Usmani, K., Valas, R., Vera, V., Villaseana, D., Waldron, L., Walker, B., Wang, J., Wang, Q., Wang, S., Warren, J., Warren, R., Wei, X., White, F., Williams, G., Willson, R., Wleczyk, R., Wooden, H., Worley, K., Wright, D., Wright, R., Wu, J., Yakub, S., Yen, J., Yoon, L., Yoon, V., Yu, F., Zhang, J., Zhou, J., Zhou, X., Zhao, S., Dunn, D., von Niederhausen, R., Weiss, R., Smith, D.R., Holt, R.A., Smith, H.O., Weinstock, G., and Gibbs, R.A.

## TITLE

JOURNAL

## REFERENCE

AUTHORS

JOURNAL

## REFERENCE

AUTHORS

JOURNAL

## COMMENT

Submitted (26-MAY-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA

3 (bases 1 to 246991)

Rat Genome Sequencing Consortium.

Submitted (20-NOV-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA

On Nov 20, 2002 this sequence version replaced gi:23666770.

The sequence in this assembly is a combination of BAC based reads and whole genome shotgun sequencing reads assembled using Atlas (http://www.hgsc.bcm.tmc.edu/projects/rat/). Each contig described in the feature table below represents a scaffold in the Atlas assembly (a 'contig-scaffold'). Within each contig-scaffold, individual sequence contigs are ordered and oriented, and separated by sized gaps filled with Ns to the estimated size. The sequence may extend beyond the ends of the clone and there may be sequence contigs within a contig-scaffold that consist entirely of whole genome shotgun sequence reads. Both end sequences and whole genome shotgun sequence only contigs will be indicated in the feature table.

----- Genome Center

Center: Baylor College of Medicine

Center code: BCM

Web site: http://www.hgsc.bcm.tmc.edu/

Contact: hgsc-help@bcm.tmc.edu

----- Project Information

Center project name: GXPC

Center clone name: CH230-17B8

----- Summary Statistics

-----

Assembly program: Phrap; version 0.990329  
 Consensus quality: 225722 bases at least Q40  
 Consensus quality: 228759 bases at least Q30  
 Consensus quality: 230490 bases at least Q20  
 Estimated insert size: 231948; sum-of-contigs estimation  
 Quality coverage: 7x in Q20 bases; sum-of-contigs estimation

-----

\* NOTE: Estimated insert size may differ from sequence length  
 (see http://www.hgsc.bcm.tmc.edu/docs/Genbank draft data.html).

\* NOTE: This is a 'working draft' sequence. It currently  
 consists of 1 contigs. Gaps between the contigs  
 are represented as runs of N. The order of the pieces  
 is believed to be correct as given, however the sizes  
 of the gaps between them are based on estimates that have  
 been provided by the submitter.

\* This sequence will be replaced  
 by the finished sequence as soon as it is available and  
 the accession number will be preserved.

\* 1 246991: contig of 246991 bp in length.

## FEATURES

source

1..246991

/organism="Rattus norvegicus"

/mol\_type="genomic DNA"

/db\_xref="taxon:10116"

/clone="CH230-17B8"

1..1165

/note="wgs contig"

complement(738413..239222)

/note="clone boundary"

clone\_end:Sp6

site:

end sequence: BH308764"

244729..246991

/note="wgs end extension"

clone\_end:Sp6"

## ORIGIN

Query Match 19.3%; Score 37; DB 14; Length 246991;  
 Best Local Similarity 54.9%; Pred. No. 1.5;  
 Matches 73; Conservative 0; Mismatches 60; Indels 0; Gaps 0;

QY 29 GACAGCATATCTTCTGATTCGTCATCTTTCATCTTCGGGGAATAGATGTGAA 88  
 |||||  
 Db 6776 GAAACCCATTTATCTCTAAGAGCTCTATGCAGATATCTAATCTTAAAGAACATATA 6717  
 |||||

QY 89 AACCTTATAAAGCGGGTTTCGAGAAACATGCTAGTATCATTTGATGACACATG 148  
 |||||  
 Db 6716 AATACCTTCAACATGTTTCTTAGTAGGATGATATCTTTTAAACCTAG 6657  
 |||||

QY 149 GACTAAGCAAAAG 161  
 |||||

Db 6656 GACTAGCGAGAGG 6644

## RESULT 8

AL671518

## LOCUS

DEFINITION Human DNA sequence from clone Rp11-315C6 on chromosome 6 Contains the 5' end of the WASF1 gene for WAS protein family member 1, the gene for pre-mRNA splicing factor 17 (PRP17, EH binding protein 3, EHB3, CDC40), a novel gene and a CpG island, complete sequence.

## ACCESSION

AL671518.3

## VERSION

GI:18855253

## KEYWORDS

HTG; CDC40; CpG island; EH binding protein; EHB3; KIAA0269;

pre-mRNA splicing factor; PRP17; SCAR1; Verprolin-homologous

protein; WAS protein family; WASF1; WAVE; WAVE1.

## SOURCE

Homo sapiens

ORGANISM

Eukaryota; Metazoa; Chordata; Vertebrata; Euteleostomi;

Mammalia; Eutheria; Euarchontoglires; Primates; Catarrhini;

Hominidae; Homo.

REFERENCE

1 (bases 1 to 67395)

AUTHORS

Tracey, A.

**TITLE**  
**JOURNAL**  
**COMMENT**

Direct Submission  
Submitted (13-MAY-2005) Wellcome Trust Sanger Institute, Hinxton, Cambridgeshire, CB10 1SA, UK. E-mail enquiries: veg@sanger.ac.uk  
Clone requests: clonerequest@sanger.ac.uk  
On Feb 21, 2002 this sequence version replaced gi:18491488.  
The following abbreviations are used to associate primary accession numbers given in the feature table with their source databases:  
Em:, EMBL; Sw:, SWISSPROT; Tr:, TREMBL; Wp:, WORMPEP; Information on the WORMPEP database can be found at  
http://www.sanger.ac.uk/Projects/C\_elegans/wormpep/ This sequence was generated from part of bacterial clone contigs of human chromosome 6, constructed by the Sanger Centre Chromosome 6 Mapping Group. Further information can be found at  
http://www.sanger.ac.uk/MGP/Chr6  
RP11-315C6 is from the library RPCI-11.2 constructed by the group of Pieter de Jong. For further details see  
http://www.chori.org/bacpac/home.htm  
VECTOR: pBACe3.6

----- Genome Center  
Center: Wellcome Trust Sanger Institute  
Center code: SC  
Web site: http://www.sanger.ac.uk  
Contact: veg@sanger.ac.uk  
-----

This sequence was finished as follows unless otherwise noted: all regions were either double-stranded or sequenced with an alternate chemistry or covered by high quality data (i.e., phred quality >= 30); an attempt was made to resolve all sequencing problems, such as compressions and repeats; all regions were covered by at least one subclone; and the assembly was confirmed by restriction digest, except on the rare occasion of the clone being a YAC.

**FEATURES****source**

```
1. 67395
/organism="Homo sapiens"
/mol_type="genomic DNA"
/db_xref="taxon:9606"
/chromosome="6"
/clone="RP11-315C6"
/clone_lib="RPCI-11.2"
2000
```

**misc\_feature**

```
join(complement(7668..7828),complement(6425..6569),
complement(4974..5013),
complement(AL590549.9:11123..11220),
complement(AL590009.10:129003..129163),
complement(AL590009.10:<114860..114994))
/genes="WASF1"
```

```
/locus_tag="RP11-181P4.1-003"
join(complement(7668..7828),complement(6425..6569),
complement(4974..5013),
complement(AL590549.9:11123..11220),
complement(AL590009.10:129003..129163),
complement(AL590009.10:<114860..114994))
/genes="WASF1"
```

**mRNA**

```
/locus_tag="RP11-181P4.1-003"
/product="WAS protein family, member 1"
/notes="match: ESTs: BG771905.1"
join(complement(7668..7828),complement(6425..6569),
complement(AL590549.9:11123..11220),
complement(AL590009.10:129003..129163),
complement(AL590009.10:114860..114994),
complement(AL590009.10:110062..110215),
complement(AL590009.10:108611..108728),
complement(AL590009.10:106941..107113),
complement(AL590009.10:106941..107113),
complement(AL590009.10:104912..105091),
complement(AL590009.10:104912..105091),
complement(AL590009.10:103122..103750),
complement(AL590009.10:101353..102213))
/genes="WASF1"
```

**gene**

```
/locus_tag="RP11-181P4.1-002"
join(complement(7668..7828),complement(6425..6569),
complement(AL590549.9:11123..11220),
complement(AL590009.10:129003..129163),
complement(AL590009.10:114860..114994),
complement(AL590009.10:106941..107113),
complement(AL590009.10:106941..107113),
complement(AL590009.10:104912..105091),
complement(AL590009.10:104912..105091),
complement(AL590009.10:103122..103750),
complement(AL590009.10:101353..102213))
/genes="WASF1"
```

**mRNA**

```
complement(AL590009.10:110062..110215),
complement(AL590009.10:108611..108728),
complement(AL590009.10:106941..107113),
complement(AL590009.10:104912..105091),
complement(AL590009.10:103122..103750),
complement(AL590009.10:101353..102213))
/genes="WASF1"
/locus_tag="RP11-181P4.1-002"
/product="WAS protein family, member 1"
/notes="match: ESTs: AA515859.1 AA885055.1 AI750432.1
AU526626.1 AL702231.1 AU142686.1 AU159926.1 AW071606.1
AW673234.1 AW956469.1 AW963370.1 BG111973.1 BG176906.1
BG665254.1 BG806840.1 BM465114.1
match: cDNAs: AF134303.1 D87459.1"
join(complement(7668..7750),complement(6425..6569),
complement(AL590009.10:129003..129163),
complement(AL590009.10:114860..114994),
complement(AL590009.10:<110062..110215))
/genes="WASF1"
/locus_tag="RP11-181P4.1-007"
join(complement(7668..7750),complement(6425..6569),
complement(AL590009.10:129003..129163),
complement(AL590009.10:114860..114994),
complement(AL590009.10:<110062..110215))
/genes="WASF1"
/product="WAS protein family, member 1"
/notes="match: ESTs: BE787088.1"
join(complement(7286..7476),complement(6425..6569),
complement(AL590549.9:11123..11220),
complement(AL590009.10:129003..129163),
complement(AL590009.10:114860..114994),
complement(AL590009.10:<110062..110215))
/genes="WASF1"
/locus_tag="RP11-181P4.1-001"
join(complement(7266..7476),complement(6425..6569),
complement(AL590009.10:129003..129163),
complement(AL590009.10:<114860..114994))
/genes="WASF1"
/locus_tag="RP11-181P4.1-006"
join(complement(7266..7476),complement(6425..6569),
complement(AL590549.9:11123..11220),
complement(AL590009.10:129003..129163),
complement(AL590009.10:114860..114994),
complement(AL590009.10:<110062..110215))
/genes="WASF1"
/locus_tag="RP11-181P4.1-001"
/product="WAS protein family, member 1"
/notes="match: ESTs: BI462105.1"
join(complement(7266..7476),complement(6425..6569),
complement(AL590009.10:129003..129163),
complement(AL590009.10:<114860..114994))
/genes="WASF1"
/locus_tag="RP11-181P4.1-006"
/product="WAS protein family, member 1"
/notes="match: ESTs: BI461486.1"
join(complement(7266..7529),
complement(AL590009.10:129003..129163),
complement(AL590009.10:114860..114994),
complement(AL590009.10:110062..110215),
complement(AL590009.10:108611..108728),
complement(AL590009.10:106941..107113),
complement(AL590009.10:106941..107113),
complement(AL590009.10:104912..105091),
complement(AL590009.10:103122..103750),
complement(AL590009.10:101353..102213))
/genes="WASF1"
/locus_tag="RP11-181P4.1-004"
join(complement(7266..7529),
complement(AL590009.10:129003..129163),
complement(AL590009.10:114860..114994),
complement(AL590009.10:110062..110215),
complement(AL590009.10:108611..108728),
complement(AL590009.10:106941..107113),
complement(AL590009.10:106941..107113),
complement(AL590009.10:104912..105091),
complement(AL590009.10:103122..103750),
complement(AL590009.10:101353..102213))
/genes="WASF1"
```

**mRNA**

```

complement(Al590009.10:104912..105091),
complement(Al590009.10:103122..103750),
complement(Al590009.10:101353..102213))
/locus tag="RP11-181P4.1-004"
/product="WAS protein family, member 1"
/notes="match: ESTs: AA85055.1 A154223.1 A1650710.1
AV664100.1 AW379048.1 AW673234.1 BF796356.1 BG111973.1
BG703130.1 BG778594.1 BG806840.1 B1544869.1 B1545690.1
B1545964.1 B1546470.1 B1550989.1 B1552818.1 B1989518.1
BM458302.1 H18215.1
match: CDNAS: AF134303.1 BC019019.1"
join(complement(7266..7497),
complement(Al590549.9:11123..11220),
complement(Al590009.10:129003..129163),
complement(Al590009.10:114860..114994),
complement(Al590009.10:110062..110215),
complement(Al590009.10:108611..108728))
/genes="WASF1"
/locus tag="RP11-181P4.1-005"
join(complement(7266..7497),
complement(Al590549.9:11123..11220),
complement(Al590009.10:129003..129163),
complement(Al590009.10:114860..114994),
complement(Al590009.10:110062..110215),
complement(Al590009.10:108611..108728))
/genes="WASF1"
/locus tag="RP11-181P4.1-005"
/product="WAS protein family, member 1"
/notes="match: ESTs: BG719866.1 B1462401.1"
join(7968..8007,8211..8460,21009..21095,29385..29514,
35333..35416,36911..37050,38534..38630,39960..40099,
40913..40987,43113..43158,45529..45630,47191..47306,
47563..47696,53994..54070,56659..56803,57781..60044)
/genes="CDC40"
/locus tag="RP1-71D21.3-005"
join(7968..8007,8211..8460,21009..21095,29385..29514,
19.0%; Score 36.4; DB 8; Length 67395;
Best Local Similarity 54.5%; Pred. No. 2.4;
Matches 73; Conservative 0; Mismatches 61; Indels 0; Gaps 0;

QY 35 CATAACTTCGTGCTGCTATTTTCATACCTTCGGGGAATAGATGTGAAACCCCT 94
|||||
Db 15983 CAAATAAATTTCAATTTTGAATATTTTAAATACCTTAAATTTGGAAGATCTCATATCCCA 16042
|||||

QY 95 TATAAACGGGGTTTCGAGAACATGCGCTAGTAGTTCATTTGACACACATGGACTAA 154
|||||
Db 16043 TATTATAAGTGGTCTAGCAACTATACACTATAAAATCTTTGGGAAGATATGAAATGTG 16102
|||||

QY 155 GCAAAAGCTGTTGT 168
|||||
Db 16103 GCTAAGCCTCATGT 16116
|||||

RESULT 9
AC154842
LOCUS
DEFINITION
Mus musculus chromosome 17 clone RP23-55C11, WORKING DRAFT
SEQUENCE, 14 unordered pieces.
AC154842 AC101665
VERSION
AC154842.1 GI:56900468
KEYWORDS
HTG; HTGS PHASE1; HTGS DRAFT; HTGS_ACTIVEPIN.
SOURCE
Mus musculus (house mouse)
ORGANISM
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Euarchontoglires; Glires; Rodentia;
Sciurognathi; Muridae; Murinae; Mus.
REFERENCE
1 (bases 1 to 317403)
Wilson,R.K.
AUTHORS
The sequence of Mus musculus clone
TITLE
Unpublished
JOURNAL
2 (bases 1 to 317403)

```

# AUTHORS TITLE JOURNAL COMMENT

Wilson,R.K.  
Direct Submission  
Submitted (30-DEC-2004) Genome Sequencing Center, 4444 Forest Park  
Parkway, St. Louis, MO 63108, USA  
On Dec 30, 2004 this sequence version replaced gi:45680671.

----- Genome Center -----  
Center: Washington University Genome Sequencing Center  
Center code: WUGSC  
Web site: http://genome.wustl.edu

Contact: submissions@wustl.edu  
----- Project Information -----  
Center project name: M\_BA0055C11  
Drafting center: WIBR

----- Summary Statistics -----  
Sequencing vector: M13; 0%  
Chemistry: Dye-primer ET; 0% of reads

Chemistry: Dye-terminator Big Dye; 100% of reads  
Assembly program: Phrap; version 0.990319  
Consensus quality: 312509 bases at least Q40  
Consensus quality: 312620 bases at least Q30  
Consensus quality: 314237 bases at least Q20

\* NOTE: This is a 'working draft' sequence. It currently  
\* consists of 14 contigs. The true order of the pieces  
\* is not known and their order in this sequence record is  
\* arbitrary. Gaps between the contigs are represented as  
\* runs of N, but the exact sizes of the gaps are unknown.  
\* This record will be updated with the finished sequence  
\* as soon as it is available and the accession number will  
\* be preserved.

\* 1 1281: contig of 1281 bp in length  
\* 1282 1381: gap of unknown length  
\* 1382 2759: contig of 1378 bp in length  
\* 2760 2859: gap of unknown length  
\* 2860 6382: contig of 3523 bp in length  
\* 6383 6482: gap of unknown length  
\* 6483 13384: contig of 6902 bp in length  
\* 13385 13485: gap of unknown length  
\* 13485 19720: contig of 6236 bp in length  
\* 19721 19820: gap of unknown length  
\* 19821 26341: contig of 6521 bp in length  
\* 26342 26441: gap of unknown length  
\* 26442 34060: contig of 7619 bp in length  
\* 34061 34150: gap of unknown length  
\* 34150 53672: contig of 19512 bp in length  
\* 53673 53772: gap of unknown length  
\* 53773 65274: contig of 11502 bp in length  
\* 65275 65374: gap of unknown length  
\* 65375 76706: contig of 11332 bp in length  
\* 76707 76806: gap of unknown length  
\* 76807 102682: contig of 25876 bp in length  
\* 102683 102782: gap of unknown length  
\* 102783 137874: contig of 35092 bp in length  
\* 137875 137974: gap of unknown length  
\* 137975 173841: contig of 35867 bp in length  
\* 173842 173941: gap of unknown length  
\* 173942 317403: contig of 143462 bp in length.

## FEATURES source

1..317403  
/organism="Mus musculus"  
/mol\_type="genomic DNA"  
/db\_xref="taxon:10090"  
/chromosome="17"  
/clone="RP23-55C11"  
1..1281  
/notes="assembly\_name:Contig41"  
1282..1381  
/estimated\_length=unknown  
1382..2759  
/notes="assembly\_name:Contig44"

## misc\_feature

## gap

## misc\_feature

```

gap          2760..2859
/estimated_length=unknown
misc_feature 2860..6382
/notes="assembly_name:Contig47"
gap          6383..6482
/estimated_length=unknown
misc_feature 6483..13384
/notes="assembly_name:Contig48"
gap          13385..13484
/estimated_length=unknown
misc_feature 13485..19720
/notes="assembly_name:Contig49"
gap          19721..19820
/estimated_length=unknown
misc_feature 19821..26341
/notes="assembly_name:Contig50"
gap          26342..26441
/estimated_length=unknown
misc_feature 26442..34060
/notes="assembly_name:Contig51"
gap          34061..34160
/estimated_length=unknown
misc_feature 34161..53672
/notes="assembly_name:Contig52"
gap          53673..53772
/estimated_length=unknown
misc_feature 53773..65274
/notes="assembly_name:Contig53"
gap          65275..65374
/estimated_length=unknown
misc_feature 65375..76706
/notes="assembly_name:Contig54"
gap          76707..76806
/estimated_length=unknown
misc_feature 76807..102682
/notes="assembly_name:Contig55"
gap          102683..102782
/estimated_length=unknown
misc_feature 102783..137874
/notes="assembly_name:Contig56"
gap          137875..137974
/estimated_length=unknown
misc_feature 137975..173841
/notes="assembly_name:Contig57"
gap          173842..173941
/estimated_length=unknown
misc_feature 173942..317403
/notes="assembly_name:Contig58"

ORIGIN
Query Match      19.0%; Score 36.4; DB 14; Length 317403;
Best Local Similarity 54.5%; Pred. No. 2.3;
Matches 73; Conservative 0; Mismatches 61; Indels 0; Gaps 0;

Qy 13 GGCATGAAGTGGCTGCACAGCATAATCTGTCTGATTCGTCTATTTCATACCTTCG 72
Db 92985 GGCATCTGGTGGCTTCACATGCAATACACTGCCTGATGATAATGTTGCCAGTATCTGC 93044
Qy 73 GGGAAATAGATGTCAAAACCCCTTATAAAACGCGGTTTTCGACAAACATCGCTAGTAT 132
Db 93045 ATGATCTGGTGTGAATACCATATGCAATAGCAGTACCTCCATTTGAAGCCATCGCTGTTAT 93104
Qy 133 CATGTGATGCAACA 146
Db 93105 CTCTGATTGCAAAA 93118

RESULT 10
AC140952 AC140952 170350 bp DNA linear PRI 26-SEP-2003
LOCUS Pan troglodytes BAC clone RP43-22112 from 7, complete sequence.
DEFINITION AC140952
ACCESSION AC140952.1 GI:28850254
VERSION

```

```

KEYWORDS
SOURCE
ORGANISM
HTG.
Pan troglodytes (chimpanzee)
Pan troglodytes
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Euarchontoglires; Primates; Catarrhini;
Hominiidae; Pan.
REFERENCE
1 (bases 1 to 170350)
AUTHORS
Shah,N., Meyer,R. and Bielicki,L.
TITLE
The sequence of Pan troglodytes BAC clone RP43-22112
JOURNAL
Unpublished (2001)
REFERENCE
2 (bases 1 to 170350)
AUTHORS
Sulston,J.E. and Wilson,R.
TITLE
Sequencing of Pan troglodytes
JOURNAL
Unpublished (2001)
REFERENCE
3 (bases 1 to 170350)
AUTHORS
Waterston,R.
TITLE
Direct Submission
JOURNAL
Submitted (05-MAR-2003) Department of Genetics, Washington
University, 4444 Forest Park Avenue, St. Louis, Missouri 63108, USA
REFERENCE
4 (bases 1 to 170350)
AUTHORS
Wilson,R.
TITLE
Direct Submission
JOURNAL
Submitted (26-SEP-2003) Department of Genetics, Washington
University, 4444 Forest Park Avenue, St. Louis, Missouri 63108, USA
COMMENT
----- Genome Center
Center: Washington University Genome Sequencing Center
Center code: WUGSC
Web site: http://genome.wustl.edu
Contact: submissions@wustl.edu
----- Summary Statistics
-----
Center project name: C_PT022112
-----

NOTICE: This sequence may not represent the entire insert of this
clone. It may be shorter because we only sequence overlapping
clone sections once, or longer because we provide a small overlap
between neighboring data submissions.

This sequence was finished as follows unless otherwise noted:
all regions were double stranded, sequenced with an alternate
chemistry, or covered by high quality data (i.e., phred quality >=
30); an attempt was made to resolve all sequencing problems, such
as compressions and repeats; all regions were covered by sequence
from more than one subclone; and the assembly was confirmed by
restriction digest.

MAPPING INFORMATION:
Mapping information for this clone was provided by Dr. Wes Warren,
Department of Genetics, Washington University, St. Louis MO. For
additional information about the map position of this sequence, see
http://genome.wustl.edu

SOURCE INFORMATION:
The RPCI-43 BAC Library has been constructed by Chung-Li Shu. DNA
was isolated from white blood cells obtained from a male chimpanzee
(Pan troglodytes, 'Clinton', Yerkes #C0471; birthdate: 6-6-80). The
clone and detailed information can be obtained from ResGen
(http://www.resgen.com) or Pieter de Jong and co-workers at
http://www.bacpac.chori.org.

NEIGHBORING SEQUENCE INFORMATION:
This sequence is the entire insert of the clone.
Location/Qualifiers
1. 170350
/organism="Pan troglodytes"
/mol_type="genomic DNA"
/db_xref="taxon:9598"
/chromosome="7"
/map="7"
/clone="RP43-22112"
/clone_lib="RPCI-43"

FEATURES
source
ORIGIN

```

```

Query Match      18.6%; Score 35.8; DB 8; Length 170350;
Best Local Similarity 63.2%; Pred. No. 3.7;
Matches 55; Conservative 0; Mismatches 32; Indels 0; Gaps 0;

QY 15 CCATGAAGTGGCTTGACACATATCTCTGATTCGTCATTTTCAATACCTTCGGG 74
    |||
Db 144944 CAGGCACTGGGTACATAGTATCATATTTTAAATTTCTTAAGTTTAAACATTTTAT 145003
    |||

QY 75 GAAATAGATGTGAAACCCCTTATAAA 101
    |||
Db 145004 ATAAGCATTTTGAAACACTTAAAAA 145030
    |||

RESULT 11
AC092873      171239 bp      DNA      linear      HTG 12-JUN-2002
LOCUS      Pan troglodytes clone RP43-22112, WORKING DRAFT SEQUENCE, 4 ordered
DEFINITION      pieces8
AC092873      AC092873.3      GI:21392490
VERSION      HTG; HTGS PHASE2; HTGS DRAFT.
KEYWORDS      Pan troglodytes (chimpanzee)
SOURCE      Pan troglodytes
ORGANISM      Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
      Mammalia; Eutheria; Euarchontoglires; Primates; Catarrhini;
      Homnidae; Pan.
REFERENCE      1 (bases 1 to 171239)
AUTHORS      Akhter,N., Antonellis,A., Avele,K., Beckstrom-Sternberg,S.M.,
      Benjamin,B., Blakeley,R.W., Bouffard,G.G., Breen,K., Brinkley,C.,
      Brooks,S., Dietrich,N.L., Granite,S., Guan,X., Gupta,J., Laric,P.,
      Haghighi,P., Hansen,N., Ho,S.-L., Idol,J.R., Karlins,E., Laric,P.,
      Lee-Lin,S.-Q., Legaspi,R., Maduro,Q.L., Maduro,V.B.,
      Margulies,E.H., Masiello,C., Maekeri,B., Mastrian,S.D.,
      McCloskey,J.C., McDowell,J., Paguirigan,C., Pearson,R.,
      Portnoy,M.E., Prasad,A., Schueler,M.G., Stantripop,S., Thomas,J.W.,
      Thomas,P.J., Touchman,J.W., Tsurgoun,C., Vogt,J.B., Walker,M.A.,
      Wetherby,K.D., Wiggins,L., Young,A., Zhang,L.-H. and Green,E.D.
      NISC Comparative Sequencing Initiative
      Unpublished
      2 (bases 1 to 171239)
      Green,E.D.
      Direct Submission
      Submitted (01-AUG-2001) NIH Intramural Sequencing Center, 8717
      Government Circle, Gaithersburg, MD 20877, USA
      3 (bases 1 to 171239)
      Green,E.D.
      Direct Submission
      Submitted (12-JUN-2002) NIH Intramural Sequencing Center, 8717
      Government Circle, Gaithersburg, MD 20877, USA
      On Jun 12, 2002 this sequence version replaced gi:15799603.
      ----- Genome Center
      Center: NIH Intramural Sequencing Center
      Center code: NISC
      Web site: http://www.nisc.nih.gov
      Contact: nisc.zoo@nhgri.nih.gov
      ----- Project Information
      Center project name: ani
      Center clone name: 022112

```

The sequence data in this record represents an 'enhanced' version of a Phase 2 submission. Specifically, the indicated order and orientation of each sequence contig has been established using one or more of the following: read-pair data from individual subclones, overlaps with neighboring clones, alignment with available reference sequence (e.g., human), and/or confirmation by PCR testing. In addition, the sequence assembly is based on at least 8X average coverage in Q20 bases and has been reviewed to rule out gross misassemblies, the low-quality ends of sequence contigs have been trimmed away, and each base is associated with a Phrap-derived quality score.

----- Summary Statistics

Sequencing vector: plasmid; n/a; 100% of reads

Chemistry: Dye-terminator Big Dye; 100% of reads  
 Assembly program: Phrap; version 0.990319  
 Consensus quality: 170538 bases at least Q40  
 Consensus quality: 170688 bases at least Q30  
 Consensus quality: 170863 bases at least Q20  
 Insert size: 152000; agarose-fp  
 Insert size: 168000; pulse-field-gel  
 Insert size: 170939; sum-of-contigs  
 Quality coverage: 30.12x in Q20 bases; agarose-fp  
 Quality coverage: 27.25x in Q20 bases; pulse-field-gel  
 Quality coverage: 26.78x in Q20 bases; sum-of-contigs

-----  
 \* NOTE: This is a 'working draft' sequence. It currently  
 \* consists of 4 contigs. Gaps between the contigs  
 \* are represented as runs of N. The order of the pieces  
 \* is believed to be correct as given, however the sizes  
 \* of the gaps between them are based on estimates that have  
 \* provided by the submitter.  
 \* This sequence will be replaced  
 \* by the finished sequence as soon as it is available and  
 \* the accession number will be preserved.

\* 82247: contig of 82247 bp in length  
 \* 82248 82347: gap of unknown length  
 \* 82348 83760: contig of 1413 bp in length  
 \* 83761 83860: gap of unknown length  
 \* 83861 168249: contig of 84389 bp in length  
 \* 168250 168349: gap of unknown length  
 \* 168350 171239: contig of 2890 bp in length.

## FEATURES

	source	1..171239	/organism="Pan troglodytes"
			/mol_type="genomic DNA"
			/db_xref="taxon:9598"
			/clone="RP43-22112"
			/clone_lib="RP43"
misc_feature	1..82247	/note="assembly_fragment"	
		clone end:SP6	
		vector side:left"	
misc_feature	1..62423	/note="clone overlaps with GenBank Accession Number AC093139 clone RP43-22A1 (center project name anh)"	
misc_feature	61398..171239	/note="clone overlaps with GenBank Accession Number AC098646 clone RP43-44G6 (center project name anj)"	
gap	82248..82347	/estimated_length=unknown	
misc_feature	82348..83760	/note="assembly_fragment"	
gap	83761..83860	/estimated_length=unknown	
misc_feature	83861..168249	/note="assembly_fragment"	
gap	168250..168349	/estimated_length=unknown	
misc_feature	168350..171239	/note="assembly_fragment"	
		clone end:T7	
		vector_side:right"	

## ORIGIN

Query Match 18.6%; Score 35.8; DB 14; Length 171239;  
 Best Local Similarity 63.2%; Pred. No. 3.7;  
 Matches 55; Conservative 0; Mismatches 32; Indels 0; Gaps 0;  
 QY 15 CCATGAAGTGGCTTGACACATATCTCTGATTCGTCATTTTCAATACCTTCGGG 74  
 |||  
 Db 145797 CCAGGCACCTGGGTACATAAGTATCATATTTTAAATTTCTTAAGTTTAAACATTTTAT 145856  
 |||  
 QY 75 GAAATAGATGTGAAACCCCTTATAAA 101  
 |||  
 Db 145857 ATAAGCATTTTGAAACACTTAAAAA 145883  
 |||







SOURCE  
ORGANISM  
Pan troglodytes (chimpanzee)  
Pan troglodytes  
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
Mammalia; Eutheria; Euarchontoglires; Primates; Catarrhini;  
Hominae; Pan.  
REFERENCE  
1 (bases 1 to 183888)  
Adams, S. and Bielicki, L.  
The sequence of Pan troglodytes BAC clone RP43-44D10  
Unpublished (2001)  
REFERENCE  
2 (bases 1 to 183888)  
Wilson, R. K.  
Direct Submission  
JOURNAL  
Submitted (01-OCT-2004) Genetics, Genome Sequencing Center, 4444  
Forest Park Parkway, St. Louis, MO 63108, USA  
REFERENCE  
3 (bases 1 to 183888)  
Wilson, R. K.  
Direct Submission  
JOURNAL  
Submitted (02-NOV-2004) Genetics, Genome Sequencing Center, 4444  
Forest Park Parkway, St. Louis, MO 63108, USA  
REFERENCE  
4 (bases 1 to 183888)  
Wilson, R. K.  
Direct Submission  
JOURNAL  
Submitted (27-APR-2005) Washington University School of Medicine,  
Genome Sequencing Center, 4444 Forest Park Parkway, St. Louis, MO  
63108, USA  
COMMENT  
On Nov 2, 2004 this sequence version replaced gi:53386473.  
----- Genome Center  
Center: Washington University Genome Sequencing Center  
Center code: WUGSC  
Web site: <http://genome.wustl.edu>  
Contact: [submissions@wustl.edu](mailto:submissions@wustl.edu)  
----- Summary Statistics  
Center project name: C\_PT044D10  
NOTICE:  
This sequence was finished as follows unless otherwise noted:  
all regions were double stranded, sequenced with an alternate  
chemistry, or covered by high quality data (i.e., phred quality >=  
30); an attempt was made to resolve all sequencing problems, such  
as compressions and repeats; all regions were covered by sequence  
from more than one subclone; and the assembly was confirmed by  
restriction digest.  
MAPPING INFORMATION:  
Mapping information for this clone was provided by Dr. Wes Warren,  
Department of Genetics, Washington University, St. Louis MO. For  
additional information about the map position of this sequence, see  
<http://genome.wustl.edu>  
SOURCE INFORMATION:  
The RPCI-43 BAC Library has been constructed by Chung-Li Shu. DNA  
was isolated from white blood cells obtained from a male chimpanzee  
(Pan troglodytes, 'Clint', Yerkes #C0471; birthdate: 6-6-80). The  
clone and detailed information can be obtained from ResGen  
(<http://www.resgen.com>) or Pieter de Jong and co-workers at  
<http://www.bacpac.chori.org>.  
This sequence is the entire insert of the clone.  
Location/Qualifiers  
1. 183888  
/organism="Pan troglodytes"  
/mol\_type="genomic DNA"  
/db\_xref="taxon:9598"  
/chromosome="7"  
/clone="RP43-44D10"  
/clone\_lib="RPCI-43"  
72798  
/note="Unresolved bases"  
FEATURES  
source  
18.6%; Score 35.8; DB 8; Length 183888;  
ORIGIN  
unsure  
Query Match

Best Local Similarity 63.2%; Pred. No. 3.7;  
Matches 55; Conservative 0; Mismatches 32; Indels 0; Gaps 0;  
QY 15 CCATGAAGTGGCTTGACACGACATAAATCTGCTGATTCGCTATTTTCAATACCTTCGGG 74  
Db 100188 CCAGGCACCTGGGTACATAGTATCATATTTTAAATTTCTCTATGTTTAAACATTTAT 100129  
QY 75 GAAATAGATGTGAAACCCCTTATAAAA 101  
Db 100128 ATAAGCATTTTGAACAACTTAAAAAA 100102  
RESULT 14  
AC149460/c  
LOCUS  
DEFINITION  
AC149460 181622 bp DNA linear HTG 08-JUL-2004  
Papio anubis clone RP41-44181, WORKING DRAFT SEQUENCE, 4 ordered  
pieces.  
AC149460  
VERSION  
AC149460.2 GI:50057884  
KEYWORDS  
HTG; HTGS PHASE2; HTGS DRAFT.  
SOURCE  
Papio anubis (olive baboon)  
ORGANISM  
Papio anubis  
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
Mammalia; Eutheria; Euarchontoglires; Primates; Catarrhini;  
Cercopithecoidea; Cercopithecinae; Papio.  
REFERENCE  
1 (bases 1 to 181622)  
Antonellis, A., Ayele, K., Benjamin, B., Blakesley, R.W.,  
Bouffard, G.G., Brinkley, C., Brooks, S., Chu, G., Coleman, B.,  
Coleman, H., Daki, N., Engle, J., Guan, X., Gupta, J., Haghighi, P.,  
Han, J., Hansen, N., Ho, S.-L., Hu, P., Hurle, B., Idol, J.R., Jones, C.,  
Karls, E., Kim, H., Kwong, P., Laric, P., Larson, S., Lee-Lin, S.-O.,  
Legaspi, R., Madden, M., Maduro, Q.L., Maduro, V.B., Margulies, E.H.,  
Masiello, C., Maskeri, B., McDowell, J., Mullikin, J.C., Paguirigan, C.,  
Park, M., Portnoy, M.E., Prasad, A., Puri, O., Reddix-Dugue, N.,  
Schandler, K., Schueler, M.G., Shah, K., Sison, C., Stantirpop, S.,  
Thomas, J.W., Thomas, P.J., Tsipouri, V., Vogt, J.L., Wetherby, K.D.,  
Young, A. and Green, E.D.  
NISC Comparative Sequencing Initiative  
Unpublished  
REFERENCE  
2 (bases 1 to 181622)  
Green, E.D.  
Direct Submission  
JOURNAL  
Submitted (03-JUN-2004) NIH Intramural Sequencing Center, 8717  
Grovemont Circle, Gaithersburg, MD 20877, USA  
REFERENCE  
3 (bases 1 to 181622)  
Green, E.D.  
Direct Submission  
JOURNAL  
Submitted (08-JUL-2004) NIH Intramural Sequencing Center, 8717  
Grovemont Circle, Gaithersburg, MD 20877, USA  
COMMENT  
On Jul 8, 2004 this sequence version replaced gi:48057351.  
----- Genome Center  
Center: NIH Intramural Sequencing Center  
Center code: NISC  
Web site: <http://www.nisc.nih.gov>  
Contact: [nisc\\_zoo@nhgri.nih.gov](mailto:nisc_zoo@nhgri.nih.gov)  
----- Project Information  
Center project name: hbl  
Center clone name: 441501  
The sequence data in this record represents an 'enhanced'  
version of a Phase 2 submission. Specifically, the indicated  
order and orientation of each sequence contig has been  
established using one or more of the following: read-pair  
data from individual subclones, overlaps with neighboring  
clones, alignment with available reference sequence (e.g.,  
human), and/or confirmation by PCR testing. In addition,  
the sequence assembly is based on at least 8x average  
coverage in Q20 bases and has been reviewed to rule out  
gross misassemblies, the low-quality ends of sequence  
contigs have been trimmed away, and each base is associated  
with a Phrap-derived quality score.  
----- Summary Statistics  
Sequencing vector: plasmid; n/a; 100% of reads

```

Chemistry: Dye-terminator Big Dye; 100% of reads
Assembly program: Phrap; version 0.990319
Consensus quality: 180653 bases at least Q40
Consensus quality: 180918 bases at least Q30
Consensus quality: 181167 bases at least Q20
Insert size: 180000; agarose-1p
Insert size: 181322; sum-of-contigs
Quality coverage: 10.99x in Q20 bases; agarose-1p
Quality coverage: 10.91x in Q20 bases; sum-of-contigs
-----
* NOTE: This is a 'working draft' sequence. It currently
* consists of 4 contigs. Gaps between the contigs
* are represented as runs of N. The order of the pieces
* is believed to be correct as given, however the sizes
* of the gaps between them are based on estimates that have
* provided by the submitter.
* This sequence will be replaced
* by the finished sequence as soon as it is available and
* the accession number will be preserved.
*
* 1 2622: contig of 2622 bp in length
* 2623 2722: gap of unknown length
* 132243 132243: contig of 129521 bp in length
* 132244 132343: gap of unknown length
* 132344 162447: contig of 30104 bp in length
* 162448 162547: gap of unknown length
* 162548 181622: contig of 19075 bp in length.
FEATURES             Location/Qualifiers
     source            1..181622
                        /organism="Papio anubis"
                        /mol_type="genomic DNA"
                        /db_xref="taxon:9555"
                        /clone="RP41-441E1"
                        /clone_lib="RP41"
     misc_feature      1..2622
                        /notes="BAC resource: http://bacpac.chori.org/"
                        /name="assembly_fragment"
                        /clone_end:SP6
                        vector_side:left"
2623..2722
                        /estimated_length=unknown
2723..132243
                        /notes="assembly_fragment"
132244..132343
                        /estimated_length=unknown
132344..162447
                        /notes="assembly_fragment"
132495..181622
                        /notes="clone overlaps with GenBank Accession Number
AC150199 clone RP41-18917 (center project name hbkt)"
162448..162547
                        /estimated_length=unknown
162548..181622
                        /notes="assembly_fragment"
                        missing T7 clone end on 3' end of insert"
ORIGIN
Query Match          18.2%; Score 35; DB 14; Length 181622;
Best Local Similarity 53.2%; Pred. No. 6.6;
Matches 74; Conservative 0; Mismatches 65; Indels 0; Gaps 0;

Qy 42 TTGCTCATTCGTCCTATTTTCAATACCTTCGGGGAATAGATGTAACACCTTATATAAA 101
      |||||
Db 73060 TTTTCTACTTTCCTCTCTTACAGCTACAGCAATGATCTGCTTAGCAGATTAAG 73001
      |||||

Qy 102 CGCGGGTTTTCGAGAACATGCGCTAGTATCATTTGATGCAACATGAGCTTAAGCAAAAG 161
      |||||
Db 73000 CCTATTTTCAGCATATAATATGCTTACCATGATTAATAATAGACTAAGCTTGTAT 72941
      |||||

Qy 162 TGGTTGTCCTTGACCCAA 180
      |||||
Db 72940 TTCTTTACTGCTGGACCAA 72922
      |||||

```

```

RESULT 15
AC116068
LOCUS
DEFINITION
Rattus norvegicus clone CH230-66D24, WORKING DRAFT SEQUENCE, 3
unordered pieces.
AC116068
AC116068.5 GI:30579201
HTG: HTGS_PHASE1; HTGS_DRAFT; HTGS_FULLTOP.
Rattus norvegicus (Norway rat)
Rattus norvegicus
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Euarchontoglires; Glires; Rodentia;
Sciurognathi; Murioidea; Muridae; Murinae; Rattus.
1 (bases 1 to 231844)
Muzny, D.Marie., Metzker, M.Lee., Abramson, S., Adams, C., Alder, J.,
Allen, C., Allen, H., Alsbrooks, S., Amin, A., Anguiano, D.,
Anyalebechi, V., Aoyagi, A., Ayodeji, M., Baca, E., Baden, H.,
Baldwin, D., Bandaranaike, D., Barber, M., Barnstead, M., Benahmed, F.,
Biswal, K., Blair, J., Blankenburg, K., Blyth, P., Brown, M.,
Bryant, N., Buhay, C., Burch, P., Burrell, K., Calderon, E.,
Cardenas, V., Carter, K., Cavazos, I., Ceasar, H., Center, A.,
Chacko, J., Chavez, D., Chen, G., Chen, R., Chen, Y., Chen, Z., Chu, J.,
Cleveland, C., Cockrell, R., Cox, C., Coyle, M., Cree, A., D'Souza, L.,
Davila, M.L., Davis, C., Davy-Carroll, L., De Anda, C., Dederich, D.,
Delgado, O., Denson, S., Deramo, C., Ding, Y., Dinh, H., Divya, K.,
Draper, H., Dugan-Rocha, S., Dunn, A., Durbin, K., Duval, B., Eaves, K.,
Egan, A., Escoto, M., Eugene, C., Evans, C.A., Falls, T., Fan, G.,
Fernandez, S., Finley, M., Flagg, N., Forbes, L., Foster, M., Foster, P.,
Fraser, C.M., Gabisi, A., Ganta, R., Garcia, A., Garner, T., Garza, M.,
Georgiev, E., Geer, K., Gill, R., Grady, M., Guerra, W., Guevara, W.,
Gunaratne, P., Haaland, W., Hamill, C., Hamilton, C., Hamilton, J.,
Harvey, Y., Havlak, P., Hawes, A., Henderson, N., Hernandez, J.,
Hernandez, R., Hines, S., Hladun, S.L., Hodgson, A., Hoques, M.,
Hollins, B., Howell, S., Hulyk, S., Hume, J., Idlebird, D., Jackson, A.,
Jackson, L., Jacob, L., Jiang, H., Johnson, B., Johnson, R., Jolivet, A.,
Karpach, S., Kelly, S., Kelly, S., Khan, Z., King, L., Kovar, C.,
Kowis, C., Kraft, C.L., Lebow, H., Levan, J., Lewis, L., Li, Z., Liu, J.,
Liu, J., Liu, W., Liu, Y., London, P., Longacre, S., Lopez, J.,
Lorensuewa, L., Loulsegged, H., Lozado, R., Lu, X., Ma, J.,
Maheshwari, M., Mahindartne, M., Mahmoud, M., Malloy, K., Mangum, A.,
Mangum, B., Mapua, P., Martin, K., Martin, R., Martinez, E.,
Mawhiney, S., McLeod, M.P., McNeill, T.Z., Meenen, E.,
Milosavljevic, A., Miner, G., Minja, E., Montemayor, J., Moore, S.,
Morgan, M., Morris, K., Morris, S., Munidasa, M., Murphy, M., Nair, L.,
Nankervis, C., Neal, D., Newton, N., Nguyen, N., Norris, S.,
Nwokeneme, O., Okwuonu, G., Olarnpunsagoon, A., Pal, S., Parks, K.,
Pasternak, S., Paul, H., Perez, A., Perez, L., Pfannkuch, C.,
Plopper, F., Poindexter, A., Popovic, D., Primus, E., Pu, L., L.,
Puazo, M., Quiroz, J., Rachlin, E., Reeves, K., Regier, M.A., Reigh, R.,
Reilly, B., Reilly, M., Ren, Y., Reuter, M., Richards, S., Riggs, F.,
Rives, C., Rodkey, T., Rojas, A., Rose, M., Rose, R., Ruiz, S.J.,
Sanders, W., Savery, G., Scherer, S., Scott, G., Shatsman, S., Shen, H.,
Shetty, J., Shvartsbeyn, A., Sisson, I., Sitter, C.D., Smajs, D.,
Sneed, A., Sodergren, E., Song, X.-Z., Sorelle, R., Sosa, J.,
Steimle, M., Strong, R., Sutton, A., Svatek, A., Taber, P., Taylor, C.,
Taylor, T., Thomas, N., Thomas, S., Tingey, A., Trejos, Z., Usmani, K.,
Valas, R., Vera, V., Villaseana, D., Waldron, L., Walker, B., Wang, J.,
Wang, Q., Wang, S., Warren, J., Warren, R., Wei, X., White, F.,
Williams, G., Willson, R., Wlarczyk, R., Woodden, H., Worley, K.,
Wright, D., Wright, R., Wu, J., Yakub, S., Yen, J., Yoon, L., Yoon, V.,
Yu, F., Zhang, J., Zhou, J., Zhou, X., Zhao, S., Dunn, D., von
Niederhausern, A., Weiss, R., Smith, D.R., Holt, R.A., Smith, H.O.,
Weinstock, G. and Gibbs, R.A.
Direct Submission
Unpublished
2 (bases 1 to 231844)
Worley, K.C.
Direct Submission
Submitted (24-MAR-2002) Human Genome Sequencing Center, Department
of Molecular and Human Genetics, Baylor College of Medicine, One
Baylor Plaza, Houston, TX 77030, USA
3 (bases 1 to 231844)
Rat Genome Sequencing Consortium.

```

TITLE  
JOURNAL

## Direct Submission

Submitted (13-MAY-2003) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA

## COMMENT

On May 13, 2003 this sequence version replaced oi:24818872. The sequence in this assembly is a combination of BAC based reads and whole genome shotgun sequencing reads assembled using Atlas (<http://www.hgsc.bcm.tmc.edu/projects/rat/>). Each contig described in the feature table below represents a scaffold in the Atlas assembly (a 'contig-scaffold'). Within each contig-scaffold, individual sequence contigs are ordered and oriented, and separated by sized gaps filled with Ns to the estimated size. The sequence may extend beyond the ends of the clone and there may be sequence contigs within a contig-scaffold that consist entirely of whole genome shotgun sequence reads. Both end sequences and whole genome shotgun sequence only contigs will be indicated in the feature table.

----- Genome Center

Center: Baylor College of Medicine

Center code: BCM

Web site: <http://www.hgsc.bcm.tmc.edu/>

Contact: hgsc-help@bcm.tmc.edu

----- Project Information

Center project name: GWC

Center clone name: CH230-66D24

----- Summary Statistics

Assembly program: Atlas 3.0;

Consensus quality: 226512 bases at least Q40

Consensus quality: 227588 bases at least Q30

Consensus quality: 228638 bases at least Q20

Estimated insert size: 22854; sum-of-contigs estimation

Quality coverage: 10x in Q20 bases; sum-of-contigs estimation

-----

\* NOTE: Estimated insert size may differ from sequence length  
 \* (see [http://www.hgsc.bcm.tmc.edu/docs/Genbank\\_draft\\_data.html](http://www.hgsc.bcm.tmc.edu/docs/Genbank_draft_data.html)).  
 \* NOTE: This is a 'working draft' sequence. It currently  
 \* consists of 3 contigs. The true order of the pieces  
 \* is not known and their order in this sequence record is  
 \* arbitrary. Gaps between the contigs are represented as  
 \* runs of N, but the exact sizes of the gaps are unknown.  
 \* This record will be updated with the finished sequence.  
 \* as soon as it is available and the accession number will  
 \* be preserved.

\* 1 229543: contig of 229543 bp in length  
 \* 229544 229643: gap of unknown length  
 \* 229644 230670: contig of 1027 bp in length  
 \* 230671 230770: gap of unknown length  
 \* 230771 231844: contig of 1074 bp in length.

## FEATURES

## source

1. .231844  
 /organism="Rattus norvegicus"  
 /mol\_type="genomic DNA"  
 /db\_xref="taxon:10116"  
 /clone="CH230-66D24"

## misc\_feature

1. .23343  
 /note="wgs\_end\_extension  
 clone\_end:T7"  
 complement(2941..3750)  
 /note="clone\_boundary  
 clone\_end:T7"

## misc\_feature

182235..225520  
 /note="clone boundary  
 clone\_end:Sp6  
 site:EcoRI

## misc\_feature

end\_sequence:BH264906"  
 227512..229543  
 /note="wgs\_end\_extension  
 clone\_end:Sp6"

## misc\_feature

229544..229643  
 /estimated\_length=unknown  
 230671..230770

## gap

## gap

/estimated\_length=unknown

## ORIGIN

Query Match 18.2%; Score 35; DB 14; Length 231844;  
 Best Local Similarity 55.3%; Pred.No. 6.5; Indels 0; Gaps 0;  
 Matches 68; Conservative 0; Mismatches 55; Indels 0; Gaps 0;  
 QY 14 GCATGAAGTGGCTTGACAGCATAATCTTTGTCTGATTCTGTTTCAATACCTTCGG 73  
 Db 9334 GCCTTTAAGTGTAAATAAGCAACACACTGCTTATTGTCTTTTACAAACAGCTTCAT 9393  
 QY 74 GGAATAGATGTCAAACCCCTTATAAACCGCGGTTTTCCAGAAACATCGGCTAGTATC 133  
 Db 9394 TGAATGTAAAGTCACATGCCATAAAATTCGGGGTTTTTCCACACTTATGAAATTCGGTGG 9453  
 QY 134 ATT 136  
 Db 9454 ATT 9456

Search completed: March 7, 2006, 12:15:05

Job time : 1788.23 secs

***This Page Blank (uspto)***

GenCore version 5.1.7  
Copyright (c) 1993 - 2006 Bioceleration Ltd.

OM nucleic - nucleic search, using sw model

Run on: March 7, 2006, 10:55:04 ; Search time 453.383 Seconds  
(without alignments)  
2822.383 Million cell updates/sec

Title: US-10-782-899-1\_COPY\_1\_192

Perfect score: 192  
Sequence: 1 gctggcgccggcccatga.....tgaccacaagaagatgcttt 192

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 4996997 seqs, 3332346308 residues

Total number of hits satisfying chosen parameters: 9993994

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

- N Geneseq\_21.\*  
1: Geneseqn1980s.\*  
2: Geneseqn1990s.\*  
3: Geneseqn2000s.\*  
4: Geneseqn2001as.\*  
5: Geneseqn2001bs.\*  
6: Geneseqn2002as.\*  
7: Geneseqn2002bs.\*  
8: Geneseqn2003as.\*  
9: Geneseqn2003bs.\*  
10: Geneseqn2003cs.\*  
11: Geneseqn2003ds.\*  
12: Geneseqn2004as.\*  
13: Geneseqn2004bs.\*  
14: Geneseqn2005s.\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	192	100.0	600	6	Abk52324 DNA encod
C 2	175.2	91.2	156638	6	Abq81850 Bifidobac
C 3	33.4	17.4	50000	3	Aaa64140 Nucleotid
C 4	32.6	17.0	278	3	Aac27526 Human sec
C 5	32.4	16.9	2717	4	Ab121958 Drosophil
C 6	32	16.7	110000	6	Continuation (16 o
C 7	32	16.7	110000	6	Continuation (3 of
C 8	31.2	16.2	18272	4	Aak69447 Human imm
C 9	31.2	16.2	18272	4	Aal06181 Human rep
C 10	31.2	16.2	18272	4	Ab198746 Human tes
C 11	31.2	16.2	26167	14	Adx83784 Human aci
C 12	30.6	15.9	466	13	Acn50973 Cotton an
C 13	30.6	15.9	549	12	Adj44041 Plant cdn
C 14	30.4	15.8	583	5	Abv51855 Human pro
C 15	30.4	15.8	1569	4	Aah52605 S. epider
C 16	30.4	15.8	3204	4	Aah54121 S. epider
C 17	30.2	15.7	515	4	Aak37564 Human bon
C 18	30.2	15.7	2000	11	Ac138062 Rice stre
C 19	30.2	15.7	110000	11	Continuation (3 of

C	20	30	15.6	1332	13	ADS47100
	21	30	15.6	7017	4	Aas32482 Bacterial
	22	30	15.6	25519	11	Acn44652 Mouse gen
C	23	29.8	15.5	10303	13	Ad167110 Novel Lac
	24	29.8	15.5	10304	13	Ad167125 Novel Lac
C	25	29.8	15.5	115829	13	Abd33448 Human can
C	26	29.6	15.4	1824	14	Aeb26996 Pinus rad
C	27	29.6	15.4	1824	14	Aeb26999 Pinus rad
	28	29.6	15.4	2000	8	Ada71889 Rice gene
	29	29.6	15.4	2000	11	Ac135459 Rice etre
	30	29.6	15.4	2510	13	Adx49899 Plant ful
	31	29.6	15.4	110000	14	ABE42401_28
	32	29.6	15.4	110000	14	ABE42737_14
C	33	29.6	15.4	200622	14	ABE39167_14
	34	29.6	15.4	260027	11	ACN44046
	35	29.4	15.3	487	9	ACH25462 Human adu
	36	29.4	15.3	768	10	ADH85446
C	37	29.4	15.3	867	13	ADs57789 Bacterial
C	38	29.4	15.3	2000	10	ACC60804 Gene sequ
C	39	29.4	15.3	2000	10	ADK62093 Disease t
C	40	29.4	15.3	2414	4	ABL22762 Drosophil
C	41	29.4	15.3	2537	4	ABL11638 Drosophil
C	42	29.4	15.3	2719	4	ABL06226 Drosophil
C	43	29.4	15.3	3612	13	ADT47758 Bacterial
	44	29.4	15.3	4088	4	ABL03916 Drosophil
C	45	29.2	15.2	836	4	Aak83773 Human imm

ALIGNMENTS

RESULT 1  
ABK52324  
ID ABK52324 standard; DNA; 600 BP.  
XX  
AC ABK52324;  
XX  
DT 13-AUG-2002 (first entry)  
XX  
DE DNA encoding cancer gene therapy associated protein.  
XX  
KW Solid cancer; anaerobic bacteria; antitumour active protein; cancer;  
KW gene therapy; gene; da.  
XX  
OS Bifidobacterium longum.  
XX  
FH Key Location/Qualifiers  
FT CDS 193..474  
FT /\*tag= a  
FT /product= "Cancer gene therapy protein"  
XX  
PN JP2002097144-A.  
XX  
PD 02-APR-2002.  
XX  
PF 21-SEP-2001; 2001JP-00290187.  
XX  
PR 21-SEP-2000; 2000JP-00287688.  
XX  
PA (AMAN/) AMANO A.  
PA (FUJI/) FUJIMORI M.  
XX  
DR WPI; 2002-448201/48.  
DR P-PSDB; AAU96807.  
XX  
PT Solid cancer therapy with anaerobic bacteria of Bifidobacterium sp. by  
PT tumor tissue specific delivery of a DNA encoding for an antitumor active  
PT protein or its precursor.  
XX  
PS Claim 10; Page 16; 21pp; Japanese.  
XX  
CC The invention describes a method of treating a solid cancer with  
CC anaerobic bacteria by site specific delivery of DNA encoding an

CC antitumour active protein or its precursor. This sequence encodes a  
 CC cancer gene therapy associated protein  
 XX  
 SQ Sequence 600 BP; 132 A; 165 C; 161 G; 142 T; 0 U; 0 Other;  
 Query Match 100.0%; Score 192; DB 6; Length 600;  
 Best Local Similarity 100.0%; Pred. No. 3.4e-57;  
 Matches 192; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
 QY 1 GCTGGCGCGCGCCATGAAGTGGCTTGACAGCATATCTTGTCTGATGCTCTATTT 60  
 DB 1 GCTGGCGCGCGCCATGAAGTGGCTTGACAGCATATCTTGTCTGATGCTCTATTT 60  
 QY 61 TCAATACCTTCGGGGAAATAGATGTGAAACCCCTTATAAACGGGGTTTTCGCAAAAC 120  
 DB 61 TCAATACCTTCGGGGAAATAGATGTGAAACCCCTTATAAACGGGGTTTTCGCAAAAC 120  
 QY 121 ATGCGCTAGTATCATTTGATGACCAACATGGAAGCAAAAGTGTCTGCCCTGACCCAA 180  
 DB 121 ATGCGCTAGTATCATTTGATGACCAACATGGAAGCAAAAGTGTCTGCCCTGACCCAA 180  
 QY 181 GAAGGATGCTTT 192  
 DB 181 GAAGGATGCTTT 192  
 RESULT 2  
 ABQ81850/c  
 ID ABQ81850 standard; DNA; 156638 BP.  
 XX  
 AC ABQ81850;  
 XX  
 DT 19-NOV-2002 (first entry)  
 XX  
 DE Bifidobacterium longum NCC2705 related nucleotide sequence SEQ ID:1106.  
 XX  
 KW Bifidobacterium longum NCC2705; Bifidobacterium; bacterial;  
 KW antidiarrheic; antibacterial; inhibitor of Salmonella; detection;  
 KW identification; lactic acid bacterium; diarrhoea; pathogenic bacteria;  
 KW rotavirus; food composition; pharmaceutical composition; gene; ds.  
 XX  
 OS Bifidobacterium longum.  
 OS Synthetic.  
 XX  
 PN EP1227152-A1.  
 XX  
 PD 31-JUL-2002.  
 XX  
 XX 30-JAN-2001; 2001EP-00102050.  
 XX  
 PR 30-JAN-2001; 2001EP-00102050.  
 XX  
 PA (NEST ) SOC PROD NESTLE SA.  
 XX  
 XX WPI; 2002-668397/72.  
 XX  
 XX Novel polynucleotide comprising Bifidobacterium genome sequence useful as  
 PT a probe or primer for detecting and/or identifying Bifidobacterium longum  
 PT in a biological sample.  
 XX  
 XX Disclosure; SEQ ID NO 1106; 80pp; English.  
 XX  
 CC The present invention describes a polynucleotide (I) comprising a  
 CC sequence of a Bifidobacterium genome selected from the nucleotide  
 CC sequences given in ABQ81842 and ABQ81843, or a sequence exhibiting at  
 CC least 90% identity or which hybridises with the sequences given in  
 CC ABQ81842 and ABQ81843. Also described is a polynucleotide (II) encoding a  
 CC fusion protein, comprising a sequence selected from 1097 sequences given  
 CC in ABP65258 to ABP63354 ligated in frame to a polynucleotide encoding a  
 CC heterologous polypeptide. (II) has antidiarrheic and antibacterial  
 CC activities, and can be used as an inhibitor of Salmonella. (I) (which is  
 CC a probe) is useful for the detection and/or identification of  
 CC Bifidobacterium longum in a biological sample. A carrier containing the

CC lactic acid bacterium Bifidobacterium longum NCC2705 (NCIM I-2618) can be  
 CC used for preventing and/or treating diarrhoea brought about by pathogenic  
 CC bacteria and/or rotavirus. The carrier is a food composition selected  
 CC from milk, yogurt, curd, cheese, fermented milks, milk based fermented  
 CC products, ice-creams, fermented cereal based products, milk based  
 CC powders, infant formula, pet food or a pharmaceutical composition  
 CC selected from tablets, liquid bacterial suspensions, dried oral  
 CC supplement, wet oral supplement, dry tube feeding or wet tube feeding.  
 CC (I) is useful in DNA arrays or chips to carry out analysis of the  
 CC expression of the Bifidobacterium gene. ABQ81844 to ABQ81850 represent  
 CC Bifidobacterium related nucleotide sequences given in the Sequence  
 CC Listing from the present invention but not mentioned further within the  
 CC specification. N.B. The sequence data for this patent is not represented  
 CC in the printed specification but is based on sequence information  
 CC supplied by the European Patent Office  
 XX  
 SQ Sequence 156638 BP; 32098 A; 46491 C; 46415 G; 31634 T; 0 U; 0 Other;  
 Query Match 91.2%; Score 175.2; DB 6; Length 156638;  
 Best Local Similarity 97.4%; Pred. No. 3.5e-50;  
 Matches 189; Conservative 0; Mismatches 3; Indels 2; Gaps 1;  
 QY 1 GCTGGCGCGCGCCATGAAGTGGCTTGACAGCATATCTTGTCTGATGCTCTATTT 60  
 DB 142164 GCTGGCGCGCGCCATGAAGTGGCTTGACAGCATATCTTGTCTGATGCTCTATTT 142105  
 QY 61 TC--AATACCTTCGGGGAAATAGATGTGAAACCCCTTATAAAACGGGGTTTTCGCAGAA 118  
 DB 142104 TCATACCCCTTCGGGGAAATAGATGTGAAACCCCTTATAAAACGGGGTTTTCGCAGAA 142045  
 QY 119 ACATGCGCTAGTATCATTTGATGACCAACATGGAAGCAAAAGTGTCTGCCCTGACCC 178  
 DB 142044 ACATGCGCTAGTATCATTTGATGACCAACATGGAAGCAAAAGTGTCTGCCCTGACCC 141985  
 QY 179 AAGAAGGATGCTTT 192  
 DB 141984 AAGAAGGATGCTTT 141971  
 RESULT 3  
 AAA64140/c  
 ID AAA64140 standard; DNA; 50000 BP.  
 XX  
 AC AAA64140;  
 XX  
 DT 20-DEC-2000 (first entry)  
 XX  
 DE Nucleotide sequence of a beta-tubulin antigen.  
 XX  
 KW Beta-tubulin antigen; inner ear protein; Meniere's disease; autoantibody;  
 KW chronic ear disease; autoimmune disease; ss.  
 XX  
 OS Homo sapiens.  
 XX  
 PN WO2000050593-A1.  
 XX  
 PD 31-AUG-2000.  
 XX  
 XX 25-FEB-2000; 2000WO-US0004795.  
 PF  
 XX 25-FEB-1999; 99US-0121549P.  
 PR  
 XX (UYTE-) UNIV TENNESSEE RES CORP.  
 PA  
 XX Yoo TU;  
 XX  
 XX WPI; 2000-558400/51.  
 DR  
 XX New beta-tubulin antigen in the membranous structure of the inner ear,  
 PT reactive with antibodies of patients with Meniere's disease, for  
 PT diagnosing Meniere's disease and distinguishing this disease from other  
 PT autoimmune ear diseases.  
 XX

PS Claim 3; Page 74-97; 115pp; English.

XX The present sequence encodes a beta-tubulin antigen. The protein is an antigen of the membranous structure of the inner ear protein, and is reactive with antibodies from patients having Meniere's disease.

CC Meniere's disease is a chronic ear disease with unknown etiology. Serum from patients suffering from this disease contain autoantibodies against a 30 kDa cochlear protein antigen. The disease is believed to be an autoimmune disease. The beta-tubulin antigen is useful as a target substance in diagnosing or detecting Meniere's disease and in distinguishing this disease from other autoimmune ear diseases

XX Sequence 50000 BP; 17281 A; 9480 C; 8791 G; 14448 T; 0 U; 0 Other;

Query Match 17.4%; Score 33.4; DB 3; Length 50000;

Best Local Similarity 52.5%; Pred. No. 3.2;

Matches 73; Conservative 0; Mismatches 66; Indels 0; Gaps 0;

QY 42 TTGCTGATCGTCTATTTTCAATACCTTCGGGGAATAGATGTGAAACCCCTTATAAA 101

DB 26376 TTTTCTACTTTGCTCTCTTACAAGCTACAAGCAATGAATGCTGCTCAGCAGATGAAG 26317

QY 102 CGGGGTTTTGCGAGAAACATGCGTAGTATCATTCATGACACATGAGTAAAG 161

DB 26316 CCTATTTTCAGCATATAATTCCTTGCCATGATTAAATAGAAATCAGCTAAGCATGAT 26257

QY 162 TGCTTGCTCCCTGACCAA 180

DB 26256 TTCTTTACTGGTGACCAA 26238

RESULT 4

AACT27526

ID AAC27526 standard; cDNA; 278 BP.

AC AAC27526;

XX

XX 06-OCT-2000 (first entry)

DE Human secreted protein 5' EST, SEQ ID NO: 31601.

XX Human; 5' EST; expressed sequence tag; secreted protein; cDNA isolation;

KW Gene therapy; chromosome mapping; ss.

XX Homo sapiens.

XX EP1033401-A2.

XX 06-SEP-2000.

XX 21-FEB-2000; 2000EP-00200610.

XX 26-FEB-1999; 99US-0122487P.

XX (GEST ) GENSET.

XX Dumas Milne Edwards J, Duclert A, Giordano J;

XX WPI; 2000-500381/45.

XX New nucleic acid that is a 5' expressed sequence tag (5' EST) for obtaining cDNAs and genomic DNAs that correspond to 5'ESTs and for diagnostic, forensic, gene therapy and chromosome mapping procedures.

XX Claim 1; SEQ ID NO 31601; 71pp + Sequence Listing; English.

XX The present sequence is one of a large number of 5' ESTs derived from mRNAs encoding secreted proteins. No ORF has yet been conclusively identified within the present sequence. The 5' ESTs were prepared from total human RNAs or polyA+ RNAs derived from 30 different tissues. EST sequences usually correspond mainly to the 3' untranslated region (UTR) of the mRNA because they are often obtained from oligo-dT primed cDNA libraries. Such ESTs are not well suited for isolating cDNA sequences

CC derived from the 5' ends of mRNAs and even in those cases where longer cDNA sequences have been obtained, the full 5' UTR is rarely included. 5' ESTs are derived from mRNAs with intact 5' ends and can therefore be used to obtain full length cDNAs and genomic DNAs. 5' ESTs are also used in diagnostic, forensic, gene therapy and chromosome mapping procedures.

CC They are used to obtain upstream regulatory sequences and to design expression and secretion vectors

XX Sequence 278 BP; 82 A; 44 C; 43 G; 108 T; 0 U; 1 Other;

Query Match 17.0%; Score 32.6; DB 3; Length 278;

Best Local Similarity 51.0%; Pred. No. 0.63; Mismatches 0; Gaps 0;

Matches 77; Conservative 0; Indels 74; Indels 0; Gaps 0;

QY 18 TGAAGTGGCTTGACAGCAATAATCTTCTCTGATTCCTTATTTTCAATACCTTCGGGGA 77

DB 99 TCATGTGGATTGTTTCTGCTCTTAGGTATTAAATGCTGTATTCAATAATTTKGGGAC 158

QY 78 ATAGATGTGAACACCCCTTATAAAGCGGGGTTTTCGACAAACATCGGCTAGTATCATTTG 137

DB 159 ATATATGTCTATTTCAATATATACAGATGTGTATCAGGCTGACATTTCCAAAGTGCACTG 218

QY 138 ATGACAAACATGCACTAAGCAAAAGTCTTGT 168

DB 219 CTGAGTTAGAGGTATATGTTTGTAAATTTT 249

RESULT 5

ABL21958/c

ID ABL21958 standard; DNA; 2717 BP.

XX ABL21958;

XX 26-MAR-2002 (first entry)

XX Drosophila melanogaster genomic polynucleotide SEQ ID NO 17347.

XX Drosophila; developmental biology; cell signalling; insecticide;

KW pharmaceutical; gene; ds.

XX Drosophila melanogaster.

XX WO200171042-A2.

XX 27-SEP-2001.

XX 23-MAR-2001; 2001WO-US009231.

XX 23-MAR-2000; 2000US-0191637P.

PR 11-JUL-2000; 2000US-00614150.

XX (PEKE ) PE CORP NY.

XX Venter JC, Adams M, Li PWD, Myers EW;

XX WPI; 2001-656860/75.

XX New isolated nucleic acid detection reagent for detecting 1000 or more genes from Drosophila and for elucidating cell signaling and cell-cell interactions.

XX Claim 1; SEQ ID NO 17347; 21pp + Sequence Listing; English.

XX The invention relates to an isolated nucleic acid detection reagent capable of detecting 1000 or more genes from Drosophila. The invention is useful in developmental biology and in elucidating cell signalling and cell-cell interactions in higher eukaryotes for the development of insecticides, therapeutics and pharmaceutical drugs. The invention discloses genomic DNA sequences (ABL16176-ABL30511), expressed DNA sequences (ABL01840-ABL16175) and the encoded proteins (ABB57737-ABB72072). The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at ftp.wipo.int/pub/published\_pct\_sequences

WP Sequence split into 5 fragments										LOCUS ABQ67195 Accession ABQ67195									
Fragment Name		Begin		End		Score 32; DB 6; Length 110000;		Best Local Similarity 50.7%; Pred. No. 14;		Matches 77; Conservative 0; Mismatches 75; Indels 0; Gaps 0;									
WP	ABQ67195_0	1	110000																
WP	ABQ67195_1	100001	210000																
WP	ABQ67195_2	200001	310000																
WP	ABQ67195_3	300001	410000																
WP	ABQ67195_4	400001	495269																
Query Match 16.7%; Score 32; DB 6; Length 110000;																			
Best Local Similarity 50.7%; Pred. No. 14;																			
Matches 77; Conservative 0; Mismatches 75; Indels 0; Gaps 0;																			
Qy	35	CATAATCTTGTCGATTGGTCTATTTTCAATACCTTCGGGGAATAGATGTGAAAAACCT	94																
Db	29177	CATGAACTCGGACACGCCTTCATTCTCATGTATTTCGGGAAGAGCCATTTTGAAAAACACT	29118																
Qy	95	TATAAAACGCGGTTTTCGAGAAACATGCGCTAGTATCATTTGATGACAAACATGGACTAA	154																
Db	29117	GATTACGCATGAATGTTGCGGAACACTGCTTCTACTTTTGCTGAAATGATTATTGCCGAT	29058																
Qy	155	GCAAAAGTCTTTGTCCCTCGACCCCAAGAAGGA	186																
Db	29057	GCTTCTGTAAGAAGATGCCAAACAAAGAAGA	29026																
RESULT 8																			
AAK69447/c																			
ID	AAK69447 standard; DNA; 18272 BP.																		
XX																			
AC	AAK69447;																		
XX																			
DT	06-NOV-2001 (first entry)																		
XX																			
DE	Human immune/haematopoietic antigen genomic sequence SEQ ID NO:24259.																		
XX																			
KW	Human; immune; haematopoietic; immune/haematopoietic antigen; cancer;																		
KW	cytostatic; gene therapy; vaccine; metastasis; ds.																		
XX																			
OS	Homo sapiens.																		
XX																			
PN	WO200157182-A2.																		
XX																			
PD	09-AUG-2001.																		
XX																			
PP	17-JAN-2001; 2001WO-US001354.																		
XX																			
XX	31-JAN-2000; 2000US-0179065P.																		
PR	04-FEB-2000; 2000US-0180628P.																		
PR	24-FEB-2000; 2000US-0184664P.																		
PR	02-MAR-2000; 2000US-0186350P.																		
PR	16-MAR-2000; 2000US-0189874P.																		
PR	17-MAR-2000; 2000US-0190076P.																		
PR	18-APR-2000; 2000US-0198123P.																		
PR	19-MAY-2000; 2000US-0205515P.																		
PR	07-JUN-2000; 2000US-0209467P.																		
PR	28-JUN-2000; 2000US-0214886P.																		
PR	30-JUN-2000; 2000US-0215135P.																		
PR	07-JUL-2000; 2000US-0216647P.																		
PR	07-JUL-2000; 2000US-0216880P.																		
PR	11-JUL-2000; 2000US-0217487P.																		
PR	11-JUL-2000; 2000US-0217496P.																		
PR	14-JUL-2000; 2000US-0218209P.																		
PR	26-JUL-2000; 2000US-0220964P.																		
PR	14-AUG-2000; 2000US-0224518P.																		
PR	14-AUG-2000; 2000US-0224519P.																		
PR	14-AUG-2000; 2000US-0225213P.																		
PR	14-AUG-2000; 2000US-0225214P.																		
PR	14-AUG-2000; 2000US-0225266P.																		
PR	14-AUG-2000; 2000US-0225267P.																		
PR	14-AUG-2000; 2000US-0225268P.																		
PR	14-AUG-2000; 2000US-0225270P.																		
PR	14-AUG-2000; 2000US-0225447P.																		



PR 14-AUG-2000; 2000US-0225757P.  
PR 14-AUG-2000; 2000US-0225758P.  
PR 14-AUG-2000; 2000US-0225759P.  
PR 18-AUG-2000; 2000US-0226279P.  
PR 22-AUG-2000; 2000US-0226681P.  
PR 22-AUG-2000; 2000US-0226868P.  
PR 22-AUG-2000; 2000US-0227182P.  
PR 23-AUG-2000; 2000US-0227009P.  
PR 30-AUG-2000; 2000US-0228924P.  
PR 01-SEP-2000; 2000US-0229287P.  
PR 01-SEP-2000; 2000US-0229343P.  
PR 01-SEP-2000; 2000US-0229344P.  
PR 01-SEP-2000; 2000US-0229345P.  
PR 05-SEP-2000; 2000US-0229509P.  
PR 05-SEP-2000; 2000US-0229513P.  
PR 06-SEP-2000; 2000US-0230437P.  
PR 06-SEP-2000; 2000US-0230438P.  
PR 08-SEP-2000; 2000US-0231142P.  
PR 08-SEP-2000; 2000US-0231143P.  
PR 08-SEP-2000; 2000US-0231144P.  
PR 08-SEP-2000; 2000US-0231413P.  
PR 08-SEP-2000; 2000US-0231414P.  
PR 08-SEP-2000; 2000US-0232080P.  
PR 08-SEP-2000; 2000US-0232081P.  
PR 12-SEP-2000; 2000US-0231968P.  
PR 14-SEP-2000; 2000US-0232397P.  
PR 14-SEP-2000; 2000US-0232398P.  
PR 14-SEP-2000; 2000US-0232399P.  
PR 14-SEP-2000; 2000US-0232400P.  
PR 14-SEP-2000; 2000US-0232401P.  
PR 14-SEP-2000; 2000US-0233063P.  
PR 14-SEP-2000; 2000US-0233064P.  
PR 14-SEP-2000; 2000US-0233065P.  
PR 21-SEP-2000; 2000US-0234223P.  
PR 21-SEP-2000; 2000US-0234274P.  
PR 25-SEP-2000; 2000US-0234997P.  
PR 25-SEP-2000; 2000US-0234998P.  
PR 26-SEP-2000; 2000US-0235484P.  
PR 27-SEP-2000; 2000US-0235834P.  
PR 27-SEP-2000; 2000US-0235836P.  
PR 29-SEP-2000; 2000US-0236327P.  
PR 29-SEP-2000; 2000US-0236367P.  
PR 29-SEP-2000; 2000US-0236368P.  
PR 29-SEP-2000; 2000US-0236369P.  
PR 29-SEP-2000; 2000US-0236370P.  
PR 02-OCT-2000; 2000US-0236802P.  
PR 02-OCT-2000; 2000US-0237037P.  
PR 02-OCT-2000; 2000US-0237038P.  
PR 02-OCT-2000; 2000US-0237039P.  
PR 13-OCT-2000; 2000US-0237040P.  
PR 13-OCT-2000; 2000US-0239335P.  
PR 20-OCT-2000; 2000US-0239337P.  
PR 20-OCT-2000; 2000US-0240960P.  
PR 20-OCT-2000; 2000US-0241121P.  
PR 20-OCT-2000; 2000US-0241785P.  
PR 20-OCT-2000; 2000US-0241786P.  
PR 20-OCT-2000; 2000US-0241787P.  
PR 20-OCT-2000; 2000US-0241788P.  
PR 20-OCT-2000; 2000US-0241808P.  
PR 20-OCT-2000; 2000US-0241809P.  
PR 20-OCT-2000; 2000US-0241826P.  
PR 01-NOV-2000; 2000US-0244617P.  
PR 08-NOV-2000; 2000US-0246474P.  
PR 08-NOV-2000; 2000US-0246475P.  
PR 08-NOV-2000; 2000US-0246476P.  
PR 08-NOV-2000; 2000US-0246477P.  
PR 08-NOV-2000; 2000US-0246478P.  
PR 08-NOV-2000; 2000US-0246523P.  
PR 08-NOV-2000; 2000US-0246524P.  
PR 08-NOV-2000; 2000US-0246525P.  
PR 08-NOV-2000; 2000US-0246526P.  
PR 08-NOV-2000; 2000US-0246527P.  
PR 08-NOV-2000; 2000US-0246528P.  
PR 08-NOV-2000; 2000US-0246532P.  
PR 08-NOV-2000; 2000US-0246533P.  
PR 08-NOV-2000; 2000US-0246609P.  
PR 08-NOV-2000; 2000US-0246610P.  
PR 08-NOV-2000; 2000US-0246611P.  
PR 08-NOV-2000; 2000US-0246613P.  
PR 17-NOV-2000; 2000US-0249207P.  
PR 17-NOV-2000; 2000US-0249208P.  
PR 17-NOV-2000; 2000US-0249209P.  
PR 17-NOV-2000; 2000US-0249210P.  
PR 17-NOV-2000; 2000US-0249211P.  
PR 17-NOV-2000; 2000US-0249212P.  
PR 17-NOV-2000; 2000US-0249213P.  
PR 17-NOV-2000; 2000US-0249214P.  
PR 17-NOV-2000; 2000US-0249215P.  
PR 17-NOV-2000; 2000US-0249216P.  
PR 17-NOV-2000; 2000US-0249217P.  
PR 17-NOV-2000; 2000US-0249218P.  
PR 17-NOV-2000; 2000US-0249244P.  
PR 17-NOV-2000; 2000US-0249245P.  
PR 17-NOV-2000; 2000US-0249264P.  
PR 17-NOV-2000; 2000US-0249265P.  
PR 17-NOV-2000; 2000US-0249297P.  
PR 17-NOV-2000; 2000US-0249299P.  
PR 17-NOV-2000; 2000US-0249300P.  
PR 01-DEC-2000; 2000US-0250160P.  
PR 01-DEC-2000; 2000US-0250391P.  
PR 05-DEC-2000; 2000US-0251030P.  
PR 05-DEC-2000; 2000US-0251988P.  
PR 05-DEC-2000; 2000US-0256719P.  
PR 06-DEC-2000; 2000US-0251479P.  
PR 08-DEC-2000; 2000US-0251856P.  
PR 08-DEC-2000; 2000US-0251868P.  
PR 08-DEC-2000; 2000US-0251869P.  
PR 08-DEC-2000; 2000US-0251989P.  
PR 11-DEC-2000; 2000US-0251990P.  
PR 05-JAN-2001; 2000US-0254097P.  
PR 05-JAN-2001; 2000US-0259678P.  
XX  
PA (HUMA-) HUMAN GENOME SCI INC.  
XX  
XX Rosen CA, Barash SC, Ruben SM;  
PI WPI; 2001-483426/52.  
XX  
DR Nucleic acids encoding human immune/hematopoietic antigen polypeptides,  
PT useful for preventing, diagnosing and/or treating cancers and metastasis.  
PT Disclosure; SEQ ID NO 24259; 3071pp + Sequence Listing; English.  
PS  
XX AAK54951 to AAK64702 encode the human immune/haematopoietic antigen (I)  
CC amino acid sequences given in AAK82170 to AAK91921. (I) have cytostatic  
CC activity, and can be used in gene therapy and vaccine production. (I)  
CC proteins and polynucleotides may be used in the prevention, diagnosis and  
CC treatment of diseases associated with inappropriate (I) expression. For  
CC example, they may be used to treat disorders associated with decreased  
CC expression by rectifying mutations or deletions in a patient's genome  
CC that affect the activity of (I) by expressing inactive proteins or to  
CC supplement the patient's own production of (I). Additionally, (I)  
CC polynucleotides may be used to produce the secreted (I), by inserting the  
CC nucleic acids into a host cell and culturing the cell to express the  
CC protein. (I) proteins and polynucleotides may be used to prevent,  
CC diagnose and treat immune/haematopoietic-related diseases, especially  
CC cancers and cancer metastases of haematopoietic-derived cells. AAK64703  
CC to AAK87694 represent human immune/haematopoietic antigen genomic  
CC sequences from the present invention. AAK54942 to AAK54950 and AAK82169  
CC represent sequences used in the exemplification of the present invention  
XX  
SQ Sequence 18272 BP; 3601 A; 5274 C; 5400 G; 3997 T; 0 U; 0 Other;  
Query Match 16.2%; Score 31.2; DB 4; Length 18272;  
Best Local Similarity 60.7%; Pred. No. 12;  
Matches 51; Conservative 0; Mismatches 33; Indels 0; Gaps 0;  
QY 71 CGGGGAATAGATGTGAAACCCTTATAAAACGGGGTTTTTCGAGAAACATGCGCTAGT 130

Db 17088 CGAGCAGAGAGATGAAAGGACAGAGATGTGGAGGTTCAGACACACTGGGCAGG 17029  
QY 131 ATCAATTGATGACAACTATGGACTAA 154  
Db 17028 AACACCATGAGAACAGGACTCA 17005  
RESULT 9  
AAL06181/c  
ID AAL06181 standard; DNA; 18272 BP.  
XX  
AC AAL06181;  
XX  
DT 21-NOV-2001 (first entry)  
XX  
XX Human reproductive system related antigen DNA SEQ ID NO: 8869.  
XX Human; reproductive system related antigen; reproductive system disorder;  
KW Human; reproductive system related antigen; cancer; gene therapy; ds.  
KW cancer; gene therapy; ds.  
XX  
OS Homo sapiens.  
XX  
PN W0200155320-A2.  
XX  
XX  
PD 02-AUG-2001.  
XX  
PF 17-JAN-2001; 2001WO-US0001339.  
XX  
XX 31-JAN-2000; 2000US-0179065P.  
PR 04-FEB-2000; 2000US-0180628P.  
PR 24-FEB-2000; 2000US-0184664P.  
PR 02-MAR-2000; 2000US-0186350P.  
PR 16-MAR-2000; 2000US-0189874P.  
PR 17-MAR-2000; 2000US-0190076P.  
PR 18-APR-2000; 2000US-0198123P.  
PR 19-MAY-2000; 2000US-0205515P.  
PR 07-JUN-2000; 2000US-0209457P.  
PR 28-JUN-2000; 2000US-0214886P.  
PR 30-JUN-2000; 2000US-0215135P.  
PR 07-JUL-2000; 2000US-0216647P.  
PR 07-JUL-2000; 2000US-0216880P.  
PR 11-JUL-2000; 2000US-0217487P.  
PR 11-JUL-2000; 2000US-0217496P.  
PR 14-JUL-2000; 2000US-0218230P.  
PR 26-JUL-2000; 2000US-0220963P.  
PR 26-JUL-2000; 2000US-0220964P.  
PR 14-AUG-2000; 2000US-0224518P.  
PR 14-AUG-2000; 2000US-0224519P.  
PR 14-AUG-2000; 2000US-0225213P.  
PR 14-AUG-2000; 2000US-0225214P.  
PR 14-AUG-2000; 2000US-0225266P.  
PR 14-AUG-2000; 2000US-0225267P.  
PR 14-AUG-2000; 2000US-0225268P.  
PR 14-AUG-2000; 2000US-0225270P.  
PR 14-AUG-2000; 2000US-0225447P.  
PR 14-AUG-2000; 2000US-0225757P.  
PR 14-AUG-2000; 2000US-0225758P.  
PR 14-AUG-2000; 2000US-0225759P.  
PR 18-AUG-2000; 2000US-0226279P.  
PR 22-AUG-2000; 2000US-0226681P.  
PR 22-AUG-2000; 2000US-0226868P.  
PR 22-AUG-2000; 2000US-0227182P.  
PR 23-AUG-2000; 2000US-0227009P.  
PR 30-AUG-2000; 2000US-0228924P.  
PR 01-SEP-2000; 2000US-0229287P.  
PR 01-SEP-2000; 2000US-0229343P.  
PR 01-SEP-2000; 2000US-0229344P.  
PR 01-SEP-2000; 2000US-0229345P.  
PR 05-SEP-2000; 2000US-0229509P.  
PR 05-SEP-2000; 2000US-0229513P.  
PR 06-SEP-2000; 2000US-0230437P.  
PR 06-SEP-2000; 2000US-0230438P.  
PR 08-SEP-2000; 2000US-0231242P.  
PR 08-SEP-2000; 2000US-0231243P.  
PR 08-SEP-2000; 2000US-0231244P.  
PR 08-SEP-2000; 2000US-0231413P.  
PR 08-SEP-2000; 2000US-0231414P.  
PR 08-SEP-2000; 2000US-0232080P.  
PR 08-SEP-2000; 2000US-0232081P.  
PR 12-SEP-2000; 2000US-0231968P.  
PR 14-SEP-2000; 2000US-0232397P.  
PR 14-SEP-2000; 2000US-0232398P.  
PR 14-SEP-2000; 2000US-0232399P.  
PR 14-SEP-2000; 2000US-0232400P.  
PR 14-SEP-2000; 2000US-0232401P.  
PR 14-SEP-2000; 2000US-0233063P.  
PR 14-SEP-2000; 2000US-0233064P.  
PR 14-SEP-2000; 2000US-0233065P.  
PR 21-SEP-2000; 2000US-0234223P.  
PR 21-SEP-2000; 2000US-0234274P.  
PR 25-SEP-2000; 2000US-0234997P.  
PR 25-SEP-2000; 2000US-0234998P.  
PR 26-SEP-2000; 2000US-0235484P.  
PR 27-SEP-2000; 2000US-0235834P.  
PR 27-SEP-2000; 2000US-0235836P.  
PR 29-SEP-2000; 2000US-0236327P.  
PR 29-SEP-2000; 2000US-0236367P.  
PR 29-SEP-2000; 2000US-0236368P.  
PR 29-SEP-2000; 2000US-0236369P.  
PR 29-SEP-2000; 2000US-0236370P.  
PR 02-OCT-2000; 2000US-0236802P.  
PR 02-OCT-2000; 2000US-0237037P.  
PR 02-OCT-2000; 2000US-0237038P.  
PR 02-OCT-2000; 2000US-0237039P.  
PR 02-OCT-2000; 2000US-0237040P.  
PR 13-OCT-2000; 2000US-0239335P.  
PR 13-OCT-2000; 2000US-0239337P.  
PR 20-OCT-2000; 2000US-0240960P.  
PR 20-OCT-2000; 2000US-0241221P.  
PR 20-OCT-2000; 2000US-0241785P.  
PR 20-OCT-2000; 2000US-0241786P.  
PR 20-OCT-2000; 2000US-0241787P.  
PR 20-OCT-2000; 2000US-0241808P.  
PR 20-OCT-2000; 2000US-0241826P.  
PR 01-NOV-2000; 2000US-0244617P.  
PR 08-NOV-2000; 2000US-0246474P.  
PR 08-NOV-2000; 2000US-0246475P.  
PR 08-NOV-2000; 2000US-0246476P.  
PR 08-NOV-2000; 2000US-0246477P.  
PR 08-NOV-2000; 2000US-0246523P.  
PR 08-NOV-2000; 2000US-0246524P.  
PR 08-NOV-2000; 2000US-0246525P.  
PR 08-NOV-2000; 2000US-0246526P.  
PR 08-NOV-2000; 2000US-0246527P.  
PR 08-NOV-2000; 2000US-0246528P.  
PR 08-NOV-2000; 2000US-0246532P.  
PR 08-NOV-2000; 2000US-0246609P.  
PR 08-NOV-2000; 2000US-0246610P.  
PR 08-NOV-2000; 2000US-0246611P.  
PR 08-NOV-2000; 2000US-0246613P.  
PR 17-NOV-2000; 2000US-0249207P.  
PR 17-NOV-2000; 2000US-0249208P.  
PR 17-NOV-2000; 2000US-0249209P.  
PR 17-NOV-2000; 2000US-0249210P.  
PR 17-NOV-2000; 2000US-0249211P.  
PR 17-NOV-2000; 2000US-0249212P.  
PR 17-NOV-2000; 2000US-0249213P.  
PR 17-NOV-2000; 2000US-0249214P.  
PR 17-NOV-2000; 2000US-0249215P.  
PR 17-NOV-2000; 2000US-0249216P.  
PR 17-NOV-2000; 2000US-0249217P.  
PR 17-NOV-2000; 2000US-0249218P.  
PR 17-NOV-2000; 2000US-0249244P.

PR 17-NOV-2000; 2000US-0249245P.  
PR 17-NOV-2000; 2000US-0249264P.  
PR 17-NOV-2000; 2000US-0249265P.  
PR 17-NOV-2000; 2000US-0249297P.  
PR 17-NOV-2000; 2000US-0249299P.  
PR 17-NOV-2000; 2000US-0249300P.  
PR 01-DEC-2000; 2000US-0250160P.  
PR 01-DEC-2000; 2000US-0250391P.  
PR 05-DEC-2000; 2000US-0251030P.  
PR 05-DEC-2000; 2000US-0251988P.  
PR 05-DEC-2000; 2000US-0256719P.  
PR 06-DEC-2000; 2000US-0251479P.  
PR 08-DEC-2000; 2000US-0251856P.  
PR 08-DEC-2000; 2000US-0251868P.  
PR 08-DEC-2000; 2000US-0251869P.  
PR 08-DEC-2000; 2000US-0251989P.  
PR 08-DEC-2000; 2000US-0251990P.  
PR 11-DEC-2000; 2000US-0254097P.  
PR 05-JAN-2001; 2001US-0259678P.  
XX  
PA (HUMA-) HUMAN GENOME SCI INC.  
XX  
XX Rosen CA, Barash SC, Ruben SM;  
XX  
XX WPI; 2001-465570/50.  
XX  
PT Isolated nucleic acid molecule encoding a reproductive system antigen is  
PT used in preventing, treating or ameliorating a medical condition.  
XX  
PS Disclosure; SEQ ID NO 8869; 1297bp + Sequence Listing; English.  
XX  
CC The present invention provides the protein and coding sequences of a  
CC number of human reproductive system related antigens. These can be used  
CC in the prevention and treatment of reproductive system disorders,  
CC including cancer. The present sequence is a genomic sequence encoding a  
CC protein of the invention  
XX  
SQ Sequence 18272 BP; 3601 A; 5274 C; 5400 G; 3997 T; 0 U; 0 Other;  
Query Match 16.2%; Score 31.2; DB 4; Length 18272;  
Best Local Similarity 60.7%; Pred. No. 12;  
Matches 51; Conservative 0; Mismatches 33; Indels 0; Gaps 0;  
QY 71 CGGGGAATAGATGTGAACACCTTATAAACCGGGGTTTTTCGAGAAACATCGCTAGT 130  
Db 17088 CGAGCAGAGAGATCAAAAGGCAGAGATGTGGAGGTGTTCAGCAGACAGCTGGGCAGG 17029  
QY 131 ATCAATTGATGACAACTGGACTAA 154  
Db 17028 AACACCCATGAGAACAGGGACTCA 17005  
RESULT 10  
ABL98746/C  
ID ABL98746 standard; DNA; 18272 BP.  
XX  
AC ABL98746;  
XX  
DT 21-JUN-2002 (first entry)  
XX  
DE Human testicular antigen encoding DNA fragment SEQ ID NO: 3398.  
XX  
KW Human; testicular antigen; testes; cancer; metastasis; immune disorder;  
KW reproductive system disorder; urinary system disorder; gene therapy;  
KW cardiovascular disorder; respiratory disorder; neurological disorder;  
KW gastrointestinal disease; infection; cytostatic; gene; ds.  
XX  
OS Homo sapiens.  
XX  
XX WO200155317-A2.  
PN  
XX  
XX 02-AUG-2001.  
PD  
XX

PF 17-JAN-2001; 2001WO-US001329.  
XX  
PR 31-JAN-2000; 2000US-0179065P.  
PR 04-FEB-2000; 2000US-0180628P.  
PR 24-FEB-2000; 2000US-0184564P.  
PR 02-MAR-2000; 2000US-0186350P.  
PR 16-MAR-2000; 2000US-0189874P.  
PR 17-MAR-2000; 2000US-0190076P.  
PR 18-APR-2000; 2000US-0198123P.  
PR 19-MAY-2000; 2000US-0205515P.  
PR 07-JUN-2000; 2000US-0209467P.  
PR 28-JUN-2000; 2000US-0214886P.  
PR 30-JUN-2000; 2000US-0215135P.  
PR 07-JUL-2000; 2000US-0216647P.  
PR 07-JUL-2000; 2000US-0216880P.  
PR 11-JUL-2000; 2000US-0217487P.  
PR 11-JUL-2000; 2000US-0217496P.  
PR 14-JUL-2000; 2000US-0218290P.  
PR 26-JUL-2000; 2000US-0220963P.  
PR 26-JUL-2000; 2000US-0220964P.  
PR 14-AUG-2000; 2000US-0224518P.  
PR 14-AUG-2000; 2000US-0224519P.  
PR 14-AUG-2000; 2000US-0225213P.  
PR 14-AUG-2000; 2000US-0225266P.  
PR 14-AUG-2000; 2000US-0225267P.  
PR 14-AUG-2000; 2000US-0225268P.  
PR 14-AUG-2000; 2000US-0225270P.  
PR 14-AUG-2000; 2000US-0225447P.  
PR 14-AUG-2000; 2000US-0225577P.  
PR 14-AUG-2000; 2000US-0225758P.  
PR 14-AUG-2000; 2000US-0225759P.  
PR 18-AUG-2000; 2000US-0226279P.  
PR 22-AUG-2000; 2000US-0226681P.  
PR 22-AUG-2000; 2000US-0226868P.  
PR 22-AUG-2000; 2000US-0227182P.  
PR 23-AUG-2000; 2000US-0227009P.  
PR 30-AUG-2000; 2000US-0228924P.  
PR 01-SEP-2000; 2000US-0229287P.  
PR 01-SEP-2000; 2000US-0229343P.  
PR 01-SEP-2000; 2000US-0229344P.  
PR 05-SEP-2000; 2000US-0229345P.  
PR 05-SEP-2000; 2000US-0229509P.  
PR 05-SEP-2000; 2000US-0229513P.  
PR 06-SEP-2000; 2000US-0230437P.  
PR 06-SEP-2000; 2000US-0230438P.  
PR 08-SEP-2000; 2000US-0231242P.  
PR 08-SEP-2000; 2000US-0231243P.  
PR 08-SEP-2000; 2000US-0231244P.  
PR 08-SEP-2000; 2000US-0231413P.  
PR 08-SEP-2000; 2000US-0231414P.  
PR 08-SEP-2000; 2000US-0232080P.  
PR 08-SEP-2000; 2000US-0232081P.  
PR 12-SEP-2000; 2000US-0231968P.  
PR 14-SEP-2000; 2000US-0232397P.  
PR 14-SEP-2000; 2000US-0232398P.  
PR 14-SEP-2000; 2000US-0232399P.  
PR 14-SEP-2000; 2000US-0232400P.  
PR 14-SEP-2000; 2000US-0232401P.  
PR 14-SEP-2000; 2000US-0232406P.  
PR 14-SEP-2000; 2000US-0233064P.  
PR 14-SEP-2000; 2000US-0233065P.  
PR 21-SEP-2000; 2000US-0234223P.  
PR 21-SEP-2000; 2000US-0234274P.  
PR 25-SEP-2000; 2000US-0234997P.  
PR 25-SEP-2000; 2000US-0234998P.  
PR 26-SEP-2000; 2000US-0235484P.  
PR 27-SEP-2000; 2000US-0235834P.  
PR 27-SEP-2000; 2000US-0235836P.  
PR 29-SEP-2000; 2000US-0236327P.  
PR 29-SEP-2000; 2000US-0236367P.  
PR 29-SEP-2000; 2000US-0236368P.  
PR 29-SEP-2000; 2000US-0236369P.



```
FT      /tag= k
FT      /number= 4
FT      7101..7197
FT      /tag= l
FT      /number= 5
FT      7198..7590
FT      /tag= m
FT      /number= 5
FT      7591..7710
FT      /tag= n
FT      /number= 6
FT      7711..7789
FT      /tag= o
FT      /number= 6
FT      7790..7908
FT      /tag= p
FT      /number= 7
FT      7909..7997
FT      /tag= q
FT      /number= 7
FT      7998..8129
FT      /tag= r
FT      /number= 8
FT      8130..9238
FT      /tag= s
FT      /number= 8
FT      9239..9349
FT      /tag= t
FT      /number= 9
FT      9350..10020
FT      /tag= u
FT      /number= 9
FT      10021..10134
FT      /tag= v
FT      /number= 10
FT      10135..10234
FT      /tag= w
FT      /number= 10
FT      10235..10319
FT      /tag= x
FT      /number= 11
FT      10320..11275
FT      /tag= y
FT      /number= 11
FT      11276..11393
FT      /tag= z
FT      /number= 12
FT      11394..11870
FT      /tag= aa
FT      /number= 12
FT      11871..12004
FT      /tag= ab
FT      /number= 13
FT      12005..12168
FT      /tag= ac
FT      /number= 13
FT      12169..12320
FT      /tag= ad
FT      /number= 14
FT      12321..12510
FT      /tag= ae
FT      /number= 14
FT      12511..12659
FT      /tag= af
FT      /number= 15
FT      12660..16257
FT      /tag= ag
FT      /number= 15
FT      13030..13308
FT      /tag= ah
FT      /note= "Alu repeat"
FT      16258..16399
FT      /tag= ai
```

```
FT      intron
FT      /number= 16
FT      16400..16907
FT      /tag= aj
FT      /number= 16
FT      16908..17057
FT      /tag= ak
FT      /number= 17
FT      17058..17504
FT      /tag= al
FT      /number= 17
FT      17505..17669
FT      /tag= am
FT      /number= 18
FT      17670..17964
FT      /tag= an
FT      /number= 18
FT      17965..18117
FT      /tag= ao
FT      /number= 19
FT      18118..18584
FT      /tag= ap
FT      /number= 19
FT      /cons splice= (5'site:NO, 3'site:YBS)
FT      18585..19194
FT      /tag= aq
FT      /number= 20
FT      18642..18644
FT      /tag= ar
FT      /note= "Translational stop codon"
FT      19172..19177
FT      /tag= as
FT      19510..19540
FT      /tag= at
FT      /note= "CT-rich region"
FT      20260..20384
FT      /tag= au
FT      /note= "Half Alu repeat"
FT      20385..20675
FT      /tag= av
FT      /note= "Alu repeat"
FT      20685..20834
FT      /tag= aw
FT      /note= "Half Alu repeat"
FT      20845..21132
FT      /tag= ax
FT      /note= "Alu repeat"
FT      22000..22320
FT      /tag= ay
FT      /note= "Alu repeat"
FT      22675..22964
FT      /tag= az
FT      /note= "Alu repeat"
FT      24649..24923
FT      /tag= ba
FT      /note= "Alu repeat"
FT      FT
FT      US6858425-B1.
FT      PN
FT      XX
FT      22-FEB-2005.
FT      PD
FT      XX
FT      03-DEC-1999; 99US-00454466.
FT      PF
FT      XX
FT      04-DEC-1998; 98US-0110850P.
FT      PR
FT      02-MAR-1999; 99US-0122550P.
FT      XX
FT      (GENZ ) GENZYME CORP.
FT      PA
FT      XX
FT      Heus JJ;
FT      PI
FT      XX
FT      WPI; 2005-170882/18.
FT      DR
FT      XX
FT      New recombinant polynucleotide comprises an alpha S1-casein promoter or
FT      an alpha glucosidase promoter, useful for designing transgenes for
```

PT	expression of human acid alpha glucosidase in the milk of transgenic animals.
PT	
XX	
PS	Disclosure; SEQ ID NO 1; 49pp; English.
XX	
CC	The invention relates to a recombinant polynucleotide comprising an alpha SI-casein promoter or an alpha glucosidase promoter operably linked to a heterologous coding sequence. The recombinant polynucleotide is useful for designing transgenes for expression of human acid alpha glucosidase in the milk of transgenic animals. It is also useful for the design of
CC	
CC	
CC	
	Query Watch Best Local Similarity 16.2%; Score 31.2; DB 14; Length 26167; Matches 51; Conservative 0; Mismatches 33; Indels 0; Gaps 0;
QY	71 CGGGGAAATAGATGTGMAAACCTTTATAAACCGCGTTTTTCGCAGAAAACATGCGCTAGT 130
Db	17958 CGAGCAGACGAGATGMAAAGCAGCATGTGTGAAGGTCTCAGCAGACAGCTGGGCAGG 17899
QY	131 ATCATTGATGACAACATGGACTAA 154
Db	17898 AACACCCTAGAGAACAGGACTCA 17875

```

RESULT 12
ACN50973
ID ACN50973 standard; cDNA; 466 BP.
XX
XX ACN50973;
XX
XX 02-DEC-2004 (first entry)
XX
XX
XX Cotton androecium tissue EST Clone ID: LIB3828-002-Q1-N6-F6, SEQ:5754.
XX
XX Cotton; plant; EST; expressed sequence tag; transgenic plant; androecium;
XX variety Nucotton378; library LIB3828; molecular tag; molecular marker;
XX genetic mapping; molecular mapping; seed germination; plant growth;
XX plant quality; plant yield; plant breeding; tissue printing; ss.
XX
XX Gossypium hirsutum.
XX OS
XX US2004123340-A1.
XX PN
XX
XX 24-JUN-2004.
XX
XX
XX 12-DEC-2001; 2001US-00021323.
XX PF
XX
XX 14-DEC-2000; 2000US-0255619P.
XX PR
XX
XX (DEIK/) DEIKMAN J.
XX PA
XX (FENG/) FENG P C C.
XX PA
XX (FINC/) FINCHER K L.
XX PA
XX (ZIEG/) ZIEGLER T E.
XX
XX Deikman J, Feng PCC, Fincher KL, Ziegler TE;
XX PI
XX WPI: 2004-479808/45.
XX DR

```

New isolated nucleic acid molecule that encodes a plant protein or its  
 fragment, useful for isolating a variety of agronomically significant  
 genes associated with plant growth, quality or yield, and as molecular  
 tags to map genes.  
 Claim 1; SEQ ID NO 5754; 34pp; English.  
 The invention relates to 17880 cotton expressed sequence tags (ESTs;  
 ACN45320-ACN63099). The ESTs were isolated from cDNA libraries generated  
 from primed or non-primed seeds from variety DP50B, mature seeds from  
 variety Coker 312 Boswell 96 Field, and androecium tissue, gynoecium  
 tissue, developing fibres, carpel walls and septa from variety  
 CCNorton33B. The invention also relates to substantially purified  
 proteins or their fragments encoded by nucleic acid molecules of the  
 invention, and to transformed plants having a nucleic acid construct

comprising a nucleic acid of the invention. The cotton ESTs are useful as molecular tags to isolate genetic regions, to isolate genes, to map genes, to determine gene function and to determining whether genes are members of a particular gene family. The nucleic acid molecules may be used for isolating a variety of agronomically significant genes associated with plant growth, quality, yield, and could also serve as links in metabolic and catabolic pathways. The nucleic acid molecules are also useful for identifying genes important in initiating and maintaining seed germination or that may be used to mitigate stresses encountered during seed germination. The ESTs additionally enable the acquisition of promoters and cis-regulatory elements which will be useful to express agronomically significant genes in these tissues and/or other tissues, and also permits the acquisition of molecular markers useful in breeding schemes, genetic and molecular mapping, and in cloning of agronomically significant genes. The nucleic acid molecules are further useful for detecting the expression level or pattern of a protein or mRNA and for detecting the presence or quantity of a protein by tissue printing. The present sequence represents a specifically claimed EST isolated from a cotton variety Nuotcon13B androecium tissue cDNA library (L183828). The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from the US patent office at [seqdata.uspto.gov/sequence.html?DocID=US20040123340](http://seqdata.uspto.gov/sequence.html?DocID=US20040123340)

	Query Match	15.9%;	Score 30.6;	DB 13;	Length 466;
	Best Local Similarity	58.1%;	Pred. No. 4;		
	Matches	54;	Conservative	0;	Mismatches 39; Indels 0; Gaps 0;
Qy	76	AAATAGATGTGAAAACCCCTTATATAAAGCGGGGTTTTTCGCAGAAACATGCGCTAGTATCAT	135		
Db	27	ATATATATCATATGTCTTTTCATTAACCTCTGGTCTCGGACTACATAATCCTTTATTATCAT	86		
Qy	136	TGATGACAAACATGGACTAAGCAAAAGTGCTTGT	168		
Db	87	TGATCATACCGGGAGGAACCAAGAGCCCTTGT	119		

RESULT 13	
ADJ44041/c	
ID	ADJ44041 standard; cDNA; 549 BP.
XX	
AC	ADJ44041;
XX	
DT	06-MAY-2004 (first entry)
XX	
DE	Plant cDNA #5041.
XX	
KW	Plant; gene; ss; transcription; plant genome augmentation; cereal;
KW	soybean; alfalfa; sunflower; canola; cotton; peanut; tobacco; sugar beet;
KW	maize; barley; sorghum; rice; wheat; crop plant; insecticide resistance;
KW	stress tolerance; salt tolerance; cold tolerance; drought tolerance;
KW	plant nutrition; apical dominance; dwarfism; early flowering; antiviral;
XX	antifungal.
XX	
OS	Rukaryota.
XX	
PN	US2004016025-A1.
XX	
PD	22-JAN-2004.
XX	
XX	26-SEP-2002; 2002US-00260238.
XX	
PR	26-SEP-2001; 2001US-0325277P.
PR	26-SEP-2001; 2001US-0325448P.
PR	04-APR-2002; 2002US-0370620P.
XX	
PA	(BUDW/) BUDWORTH P.
PA	(MOUG/) MOUGHAMER T.
PA	(BRIG/) BRIGGS S P.
PA	(COOP/) COOPER B.
PA	(GLAZ/) GLAZEBROOK J.
PA	(GOFF/) GOFF S A.

PA (KATA/) KATAGIRI F.  
PA (KREP/) KREPS J.  
PA (PROV/) PROVART N.  
PA (RICK/) RIQUE D.  
PA (ZHUT/) ZHU T.  
XX  
XX Budworth P, Moughamer T, Briggs SP, Cooper B, Glazebrook J;  
PI Goff SA, Katagiri F, Kreps J, Provart N, Rique D, Zhu T;  
XX  
XX WPI; 2004-190374/18.  
XX  
XX New rice promoter, useful for manipulating crop plants to alter or  
PT improve phenotypic characteristics, e.g. produce large quantities of oil  
PT or proteins, resistance to insecticides, virus or fungi, stress tolerance  
PT or high nutritional value.  
XX  
XX Example 13; SEQ ID NO 5041; 230pp; English.  
XX  
XX The invention relates to plant nucleotide sequences that direct seed-,  
CC leaf- and/or stem-, panicle-, root- or pollen-specific or -preferential  
CC or constitutive transcription of an operatively linked nucleic acid  
CC segment. The invention also relates to a method for augmenting a plant  
CC genome and a method of identifying a gene, where its expression is  
CC altered in the seed, leaf, stem, panicle, pollen, root or is constitutive  
CC in a plant cell. The plant is a cereal, e.g. soybean, alfalfa, sunflower,  
CC canola, cotton, peanut, tobacco or sugar beet, preferably maize, barley,  
CC sorghum, rice or wheat. The polynucleotides and the polypeptides they  
CC encode are useful for manipulating crop plants to alter or improve  
CC phenotypic characteristics, to produce large quantities of oil or  
CC proteins, to incur resistance to insecticides, viruses or fungi, and to  
CC incur stress tolerance (e.g. salt, cold or drought) to ensure the plants  
CC have a high nutritional value with reduced apical dominance or dwarfism,  
CC early flowering or altered metabolic pathways. This sequence represents a  
CC plant nucleic acid of the invention. Note: The sequence data for this  
CC patent did not form part of the printed specification but was obtained in  
CC electronic format directly from USPTO at [seqdata.uspto.gov/sequence.html](http://seqdata.uspto.gov/sequence.html).  
XX  
SQ Sequence 549 BP; 136 A; 143 C; 88 G; 177 T; 0 U; 5 Other;  
  
Query Match 15.9%; Score 30.6; DB 12; Length 549;  
Best Local Similarity 51.9%; Pred. No. 4.3;  
Matches 69; Conservative 0; Mismatches 64; Indels 0; Gaps 0;  
  
QY 48 GATTCTCTATTTCATACCTTCGGGAAATAGATGTGAAACCCCTTATAAAACCGGG 107  
Db 235 GAATCATTTTGAGTTCAAGACTGCTGGAAGAAAAGTTATCGAAGTCTTGACCAAGCAGTG 176  
  
QY 108 TTTTCGAGAACATCGGCTAGTATCATTTGATGACAAACATGAGCTAAGCAAAAGTCTTG 167  
Db 175 ATGACCATGAATAAGGAGAGATTTGCTTTGGTGACAAATTCACCTGAATATGCTCTTTG 116  
  
QY 168 TCCCTCGACCCAA 180  
Db 115 TCCACTGAATCAA 103  
  
RESULT 14  
ABV51855  
ID ABV51855 standard; cDNA; 583 BP.  
XX  
XX AC ABV51855;  
XX  
XX DT 17-SEP-2002 (first entry)  
XX  
XX Human prostate expression marker cDNA 51846.  
XX  
XX Human, prostate cancer; cytostatic; carcinogen; pharmacodynamic marker;  
KW pharmacogenomic marker; gene; ss.  
XX  
XX OS Homo sapiens.  
XX  
XX PN WO200160860-A2.  
XX

PD 23-AUG-2001.  
XX  
XX 20-FEB-2001; 2001WO-US005171.  
XX  
XX 17-FEB-2000; 2000US-0183319P.  
PR 16-MAR-2000; 2000US-0189862P.  
PR 25-MAY-2000; 2000US-0207454P.  
PR 09-JUN-2000; 2000US-0211314P.  
PR 18-JUL-2000; 2000US-0219007P.  
PR 13-DEC-2000; 2000US-0255281P.  
XX  
XX (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.  
XX  
XX Schlegel R, Endege WO, Monahan JE;  
XX WPI; 2001-662795/76.  
XX  
XX Novel isolated nucleic acid molecule associated with cancerous state of  
PT prostate cells and correlating with presence of prostate cancer, useful  
PT for detecting presence of prostate cancer, stage of prostate cancer.  
XX  
XX Claim 1; Page 10067-10068; 11750pp; English.  
XX  
XX The invention relates to an isolated nucleic acid molecule (I) comprising  
CC a nucleotide sequence given in Tables 1-9 (ABV00010-ABV62213) of the  
CC specification or its complement. (i) is useful for: (a) assessing whether  
CC a patient is afflicted with prostate cancer; (b) monitoring the  
CC progression of prostate cancer in a patient; (c) assessing the efficacy  
CC of a test compound to inhibit prostate cancer in a patient; (d) assessing  
CC the efficacy of a therapy for inhibiting prostate cancer in a patient;  
CC (e) selecting a composition for inhibiting prostate cancer in a patient;  
CC (f) assessing the prostate cell carcinogenic potential of a compound; (g)  
CC determining whether prostate cancer has metastasized in a patient; (h)  
CC assessing the aggressiveness or indolence of prostate cancer in a patient  
CC ; (I) is also useful as a pharmacodynamic or pharmacogenomic marker  
XX  
SQ Sequence 583 BP; 210 A; 79 C; 104 G; 188 T; 0 U; 2 Other;  
  
Query Match 15.8%; Score 30.4; DB 5; Length 583;  
Best Local Similarity 52.9%; Pred. No. 5.2;  
Matches 64; Conservative 0; Mismatches 57; Indels 0; Gaps 0;  
  
QY 49 ATTGCTATTTCATACCTTCGGGAAATAGATGTGAAACCCCTTATAAAACGGGGT 108  
Db 18 AGTCTTCTAGNATTAAATTAACCGGGATTGAGTTCATAAAAGCTATATTCTCATGTGT 77  
  
QY 109 TTTTCGAGAACATCGGCTAGTATCATTTGATGACACATGAGCTAGCAAAAGTCTTGT 168  
Db 78 TTGGGCGAGAAATTTTCAAGAAATATCATATTTCAGATTTCAGGGAAGAACTCAAAAGTTTAT 137  
  
QY 169 C 169  
Db 138 C 138  
  
RESULT 15  
AAH52605  
ID AAH52605 standard; DNA; 1569 BP.  
XX  
XX AC AAH52605;  
XX  
XX DT 03-SEP-2001 (first entry)  
XX  
XX S. epidermidis open reading frame nucleotide sequence SEQ ID NO:603.  
XX  
XX Staphylococcus epidermidis SRI strain; infection; diagnosis; vaccination;  
KW endocarditis; ds.  
XX  
XX OS Staphylococcus epidermidis.  
XX  
XX PN WO200134809-A2.  
XX  
XX PD 17-MAY-2001.

XX 09-NOV-2000; 2000NO-US030782.  
PF  
XX  
XX 09-NOV-1999; 99US-0164258P.  
PR  
XX  
XX (GLAX ) GLAXO GROUP LTD.  
PA  
XX  
XX Kimmerly WJ;  
PI  
XX  
XX WPI; 2001-316495/33.  
DR  
XX P-PSDB; AAG81755.  
PT  
XX Nucleic acids encoding polypeptides from Staphylococcus epidermidis,  
PT useful for vaccinating against infections, e.g. endocarditis.  
XX  
XX Claim 8; Page 195; 2188pp; English.  
PS  
XX  
XX AAH52304 to AAH53970 represent nucleic acids (I) encoding polypeptides  
CC (II), given in AAG81454 to AAG83120, from Staphylococcus epidermidis. (I)  
CC and (II) can have antibacterial activity and therefore can be used in  
CC vaccination. The nucleic acids (I) may be used to produce the S.  
CC epidermidis polypeptides (II) via the production of vectors containing  
CC them which are used to produce hosts cells which express the  
CC polypeptides. The polypeptides (II) (and/or nucleic acids) may then be  
CC used to vaccinate subjects and to raise antibodies against the bacteria.  
CC The polypeptides may also be used to assay for other inhibitors of their  
CC activity and therefore identify compounds that may be used for the  
CC treatment of S. epidermidis infections, e.g. endocarditis. AAH53971 to  
CC AAH55090 represent specifically claimed S. epidermidis genomic DNA  
CC polynucleotide sequences from the present invention. AAH55091 to AAH55098  
CC represent oligonucleotide sequences and primers which are used in the  
CC exemplification of the present invention. N.B. The present invention  
CC specifically claims all the polynucleotide sequences given in the  
CC sequence listing of the present specification, however the sequence  
CC listing only goes up to SEQ ID NO:4454 so even though sequences are given  
CC in the disclosure for SEQ ID NO:4465 to 4472, no sequences are present  
CC for SEQ ID NO:4455 to 4464  
XX  
XX Sequence 1569 BP; 616 A; 186 C; 274 G; 493 T; 0 U; 0 Other;

Query Match 15.8%; Score 30.4; DB 4; Length 1569;  
Best Local Similarity 57.3%; Pred. No. 8.1;  
Matches 55; Conservative 0; Mismatches 41; Indels 0; Gaps 0;  
QY 49 ATTGCTCTATTTTCATACCTTCGGGGAATAGATGTGAACCCCTTATAAAACGGCGGT 108  
DB 87 ATATGAATATGAACAAATAATAGCGGATCCATATCTCTCACTGCTTATAAACTAATGT 146  
QY 109 TTTCGCAGAAACATGCGCTAGTATCATTTGATGACAA 144  
DB 147 TAACGGGATATATTGATAGTTGATTGATGAATA 182

Search completed: March 7, 2006, 12:31:05  
Job time : 457.383 secs



Result No.	Score	Query		Length	DB	ID	Description
		Match	↑				
1	40.8	21.2	673	9	CE144436	CE144436	tigr-gss-
C 2	36.8	19.2	826	10	DU019646	DU019646	276081 TC
C 3	36.4	19.0	749	9	BH039236	BH039236	RPCI-24-2
4	35	18.2	1275	8	DN671770	DN671770	CFW64-F08
5	34.6	18.0	852	2	BG542199	BG542199	602571637
6	34.4	17.9	590	9	BH378221	BH378221	AG-ND-119
C 7	34.2	17.8	600	9	AZ964876	AZ964876	2M0234D24
8	34.2	17.8	619	3	BP519798	BP519798	BP519798
9	34.2	17.8	651	8	DN244174	DN244174	ACAAE-aaa2
10	34.2	17.8	878	7	CR281388	CR281388	CR281388
C 11	33.8	17.6	945	11	CNS0757U	AL435288	T3 end of
C 12	33.6	17.5	845	11	CNS0757U	AL435288	T3 end of
C 13	33.4	17.4	463	10	CW393812	AL439760	clone BAC
14	33.4	17.4	494	6	CA394392	CW393812	fabb001f6
15	33.4	17.4	728	9	BH085133	CA394392	CS1a07.y
16	33.2	17.3	449	7	CG1673288	BH085133	hwg1c11.b
17	33.2	17.3	467	6	CP166270	A0932B01-	A0932B01-
18	33.2	17.3	493	7	CK391130	CP166270	B0768501-
19	33.2	17.3	501	6	CD552026	CK391130	K0827F01-
20	33.2	17.3	501	6	CF1652336	CD552026	B0337C08-
21	33.2	17.3	503	6	CF168303	CF1652336	B0753H06-
22	33.2	17.3	538	6	CF162296	CF168303	B0797D04-
23	33.2	17.3	538	6	CF162296	CF162296	B0709D06-

Db 326 TGGCTTGTCTTCATTATGCTCAGCTCTCTCTGTTTTCAATAGAGTTTGTCAAATAAG 385  
 QY 83 TGTGAAAACCTTATTAACCGCGGTTTTTCGCGAGAAACATGCGCTAGTATCATTTGATGAC 142  
 Db 386 TAGCAAGAGTTTGATAGCAGGAGGTTTTAGCTTAGTCTGGCTATCGGTTCACTGACTGT 445  
 QY 143 AACATCGACTAA 154  
 Db 446 ATTCTGTAGTAA 457

RESULT 2  
 DU019646/c  
 LOCUS  
 DEFINITION 276061 Tomato MboI BAC Library lycopersicon esculentum genomic  
 DU019646 clone SL MboI0087H18 5, genomic survey sequence.  
 VERSION DU019646.1 GI:72421548  
 KEYWORDS GSS.  
 SOURCE Lycopersicon esculentum (Solanum lycopersicum)  
 ORGANISM Lycopersicon esculentum  
 Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;  
 Spermatophyta; Magnoliophyta; eudicotyledons; core eudicotyledons;  
 asterids; lamiids; Solanales; Solanaceae; Solanum; Lycopersicon.  
 REFERENCE 1 (bases 1 to 826)  
 AUTHORS Mueller L.A., Buel, R.M., Wang, Y., Tanksley, S.D., Giovannoni, J.J.,  
 Van Eck, J. and Stack, S.  
 TITLE BAC end sequencing from three Solanum lycopersicon libraries  
 JOURNAL Unpublished (2005)  
 COMMENT Other GSSs: 286317  
 Contact: Lukas Mueller  
 Tankley Lab, Dept. of Plant Breeding  
 Cornell University  
 251 Emerson Hall, Ithaca, NY 14853, USA  
 Tel: 607-255-6557  
 Fax: 607-255-6683  
 Email: sgn-feedback@sgn.cornell.edu  
 Plate: 87 row: H column: 18  
 Seq primer: T7  
 Class: BAC ends  
 High quality sequence start: 16  
 High quality sequence stop: 613.

FEATURES  
 source  
 1..826  
 /organism="Lycopersicon esculentum"  
 /mol\_type="genomic DNA"  
 /cultivar="Heinz 1706"  
 /db\_xref="taxon:4081"  
 /clone="SL MboI0087H18"  
 /lab\_host="E. coli"  
 /clone\_lib="Tomato MboI BAC Library"  
 /notes="Vector: pBelobAC11; Site\_1: MboI"

ORIGIN  
 Query Match 19.2%; Score 36.8; DB 10; Length 826;  
 Best Local Similarity 59.6%; Pred. No. 1.9; Indels 0; Gaps 0;  
 Matches 62; Conservative 0; Mismatches 42  
 QY 70 TCGGGGAAATAGATGTGAAACCCCTTATAAACCGCGGTTTTTCGAGAAACATGCGCTAG 129  
 Db 205 TCTGGCTACTCAATGTGTAAACCTTTTATTTCTCTGGATTTCTGACTAGGAACCTTCG 146  
 QY 130 TATCATTTGATGACACATGAGCTAAGCAAAAGTCCTTGCCCT 173  
 Db 145 ATTAATTGATGAAATATGAAGTGAAGAAATAATTAATTGTCCTCT 102

RESULT 3  
 BH039236/c  
 LOCUS  
 DEFINITION RPCI-24-254N24.TV RPCI-24 Mus musculus genomic clone  
 RPCI-24-254N24, genomic survey sequence.  
 ACCESSION BH039236

VERSION  
 KEYWORDS  
 SOURCE  
 ORGANISM

REFERENCE  
 AUTHORS  
 TITLE  
 JOURNAL  
 COMMENT

BH039236.1 GI:14816964  
 GSS.  
 Mus musculus (house mouse)  
 Mus musculus  
 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
 Mammalia; Theria; Euarchontoglires; Glires; Rodentia;  
 Sciurognathi; Muridea; Muridae; Murinae; Mus.  
 1 (bases 1 to 749)  
 Zhao, S., Niernan, W., Malek, J., Shatsman, S., Akinret, B., Levine, M.,  
 Tsedaye, G., Geir, K., Krol, M., Shvartsbeyn, A., Gebregeorgis, E.,  
 Russell, D., de Jong, P. and Fraser, C.M.  
 Mouse BAC End Sequences from Library RPCI-24  
 Unpublished (1999)  
 Other GSSs: RPCI-24-254N24.TJ  
 Contact: Shaying Zhao  
 Department of Eukaryotic Genomics  
 The Institute for Genomic Research  
 9712 Medical Center Dr., Rockville, MD 20850, USA  
 Tel: 301 838 0200  
 Fax: 301 838 0208  
 Email: szhao@tigr.org

Clones are derived from the mouse BAC library RPCI-24. For BAC  
 library availability, please contact Pieter de Jong  
 (pdejong@mail.cho.org). Clones may be purchased from BACPAC  
 Resources (<http://www.choi.org/bacpac/orderingframe.htm>). BAC end  
 page: [http://www.tigr.org/tdb/bac\\_ends/mouse/bac\\_end\\_intro.html](http://www.tigr.org/tdb/bac_ends/mouse/bac_end_intro.html)  
 Plate: 254 row: N column: 24  
 Seq primer: T7  
 Class: BAC ends.

FEATURES  
 source

Location/Qualifiers  
 1..749  
 /organism="Mus musculus"  
 /mol\_type="genomic DNA"  
 /strain="C57BL/6J"  
 /db\_xref="taxon:10090"  
 /clone="RPCI-24-254N24"  
 /sex="Male"  
 /cell\_type="Spleen/Brain"  
 /clone\_lib="RPCI-24"  
 /notes="Vector: PTARBAC1; Site\_1: BamHI; Site\_2: BamHI;  
 RPCI-24 Mouse BAC Library produced by Pieter de Jong. The  
 library was cloned in the pTARBAC1 cloning vector at the  
 BamHI sites using MboI partially digested male C57BL/6J  
 DNA."

ORIGIN

Query Match 19.0%; Score 36.4; DB 9; Length 749;  
 Best Local Similarity 54.5%; Pred. No. 2.5;  
 Matches 73; Conservative 0; Mismatches 61; Indels 0; Gaps 0;  
 QY 13 GGCCATGAGTGGCTTGACAAGCATATCTTGCTGCTGCTATTTTCAATACCTTCG 72  
 Db 456 GGCAATTCGTGGCTTCACATGCATCAATCACTGCTGATGATATGTTGCCAGTATCTTGC 397  
 QY 73 GGGAAATAGATGTGAAACCCCTTATAAACCGCGGTTTTTCGAGAAACATGCGCTAGTAT 132  
 Db 396 ATGCATTTGGTGTGATACCATTAATGCGTAGCCCTCCATTTTGAAGCCATCGCTGTAT 337  
 QY 133 CATTTGATGACACA 146  
 Db 336 CTCTGATTGCAAAA 323

RESULT 4  
 DN671770  
 LOCUS  
 DEFINITION DN671770 1275 bp mRNA linear EST 29-MAR-2005  
 CFW64-F08.xid-t SHGC-CFW Gasterosteus aculeatus cDNA clone  
 CFW64-F08 3', mRNA sequence.  
 ACCESSION DN671770  
 VERSION DN671770.1 GI:61991827  
 KEYWORDS EST.  
 SOURCE Gasterosteus aculeatus (three spined stickleback)  
 ORGANISM Gasterosteus aculeatus

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Actinopterygii; Neopterygii; Teleostei; Euteleostei; Neoteleostei; Acanthomorpha; Acanthopterygii; Percomorpha; Gasterosteiformes; Gasterosteidae; Gasterosteus.

#### REFERENCE

1 (bases 1 to 1275)

#### AUTHORS

Kingsley,D.M., Peichel,C., Balabhadra,S., Grimwood,J., Dickson,M., Schmutz,J., and Myers,R.M.

#### TITLE

Expressed sequence tags from *Gasterosteus aculeatus*

#### JOURNAL

Unpublished (2003)

#### COMMENT

Contact: Grimwood, Jane

Stanford Human Genome Center

Stanford University School of Medicine

975 S California Ave, Palo Alto, CA 94304, USA

Tel: 650 320 5917

Fax: 650 320 5801

Email: jane@shgc.stanford.edu

Plate: 64

High quality sequence start: 3

High quality sequence stop: 841.

#### FEATURES

Location/Qualifiers

1..1275

/organism="Gasterosteus aculeatus"

/mol\_type="mRNA"

/strain="Conner Creek sticklebacks, WA USA"

/db\_xref="taxon:69293"

/clone="CFW64-F08"

/sex="mixed male and female"

/tissue\_type="gills"

/dev\_stage="adult"

/lab\_host="DH10B (T1 phage resistant)"

/clone\_lib="SHGC-CFW"

/notes="Vector: Express 1; Total and poly A+ RNA was isolated from the indicated stickleback tissue, and a cDNA library was constructed in the Express 1 plasmid vector by Open Biosystems. First strand cDNA synthesis was primed with an 54 bp linker primer containing an oligodT sequence preceded by a synthetic NotI site (first strand primer: 5'-GACTAGTCTAGATCGAGCGGCCCC(T)25-3'). Following second strand synthesis, cDNAs were made blunt at the end corresponding to the original 5 prime end of mRNA, and cloned directionally into the NotI and EcoRV sites of Express 1. Note that the EcoRV site is typically destroyed in the blunt end cloning, leaving a junction of the form 'xxxATC'(where ATC is the second half of the EcoRV site, and xxx is derived from the cDNA sequence). A map of the Express 1 vector is available at: [http://www.openbiosystems.com/cdna\\_library\\_construction\\_fa\\_q.php#8](http://www.openbiosystems.com/cdna_library_construction_fa_q.php#8) The primary library was transformed and amplified in DH10B (T1 phage resistant) bacteria. Clones available from Open Biosystems: <http://www.openbiosystems.com/stickleback>"

#### ORIGIN

Query Match 18.2%; Score 35; DB 8; Length 1275;

Best Local Similarity 66.7%; Pred. No. 7.9;

Matches 50; Conservative 0; Mismatches 25; Indels 0; Gaps 0;

34 GCATAATCTGTCTGATTCGTCTATTTCATACCTTCGGGGAAATAGATGTCAAACCC 93

1087 GCGACACATCTCTGAAAGTGGAAATTCGGAACCTTCGGGGAAAGGCGCGAANAACA 1146

94 TTATAAAACCGGGT 108

1147 GTTTAAACCCGGT 1161

#### RESULT 5

BG542199

LOCUS

DEFINITION

BG542199

602571637F1 NIH\_MGC\_77 Homo sapiens cDNA clone IMAGE:4696041 5', mRNA sequence.

ACCESSION

BG542199

1 GI:13534432

#### KEYWORDS

SOURCE

ORGANISM

EST.

Homo sapiens (human)

Homo sapiens

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Euarchontoglires; Primates; Catarrhini; Homnidae; Homo.

1 (bases 1 to 852)

NIH-MGC <http://mgc.nci.nih.gov/>.

National Institutes of Health, Mammalian Gene Collection (MGC)

Unpublished (1999)

JOURNAL

COMMENT

Contact: Robert Strausberg, Ph.D.

Email: [cgabbs-r@mail.nih.gov](mailto:cgabbs-r@mail.nih.gov)

Tissue Procurement: CLONTECH Laboratories, Inc.

cDNA Library Preparation: CLONTECH Laboratories, Inc.

cDNA Library Arrayed by: The I.M.A.G.E. Consortium (LLNL)

DNA Sequencing by: Incyte Genomics, Inc.

Clone distribution: MGC clone distribution information can be found through the I.M.A.G.E. Consortium/LLNL at: <http://image.llnl.gov>

Plate: LLC1522 row: j column: 10

High quality sequence stop: 365.

#### FEATURES

Location/Qualifiers

1..852

/organism="Homo sapiens"

/mol\_type="mRNA"

/db\_xref="taxon:9606"

/clone="IMAGE:4696041"

/lab\_host="DH10B (T1 phage-resistant)"

/clone\_lib="NIH\_MGC\_77"

/notes="Organ: lung; Vector: pDNR-LIB (Clontech); Site 1: SfiI (ggcgctcgcc); Site 2: SfiI (ggcattatggcc); 5' and 3' adaptors were used in cloning as follows: 5' adaptor sequence: 5'-CAGCGCATATGATGCG-3' and 3' adaptor sequence: 5'-ATTCTAGAGCGCGGCGGCGCATG-3' (where B = A, C, or G and N = A, C, G, or T). Average insert size 1.9 kb (range 0.5-4.0 kb). 12/15 colonies contained inserts by PCR. This library was enriched for full-length clones and was constructed by Clontech Laboratories (Palo Alto, CA). Note: this is a NIH\_MGC Library."

#### ORIGIN

Query Match 18.0%; Score 34.6; DB 2; Length 852;

Best Local Similarity 55.4%; Pred. No. 9.7;

Matches 67; Conservative 0; Mismatches 54; Indels 0; Gaps 0;

38 AATCTGTCTGCTGCTCTATTTCATACCTTCGGGGAAATAGATGTGAAACCCCTAT 97

446 ACTATGCTCTCTGTCGCAACGCCACATACATCGGGACCTTAAGTACCAGCCCATAT 505

98 AAAACCGCGGGTTTCGAGAAACATCGCTAGTATCATTTGATGACACATGGACTAAGCA 157

506 CAACGCGGGCTCTATCTCTACTCGGACAGTCACATTTTCCATACATGGACTACCCC 565

158 A 158

566 A 566

#### RESULT 6

BH378221

LOCUS

DEFINITION

BH378221

AG-ND-119J16.TR ND-TAM Anopheles gambiae genomic clone

AG-ND-119J16, genomic survey sequence.

ACCESSION

BH378221

VERSION

BH378221.1 GI:17324363

GSS.

KEYWORDS

ORGANISM

Anopheles gambiae (African malaria mosquito)

Anopheles gambiae

Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota; Neoptera; Endopterygota; Diptera; Nematocera; Culicoidea; Culicidae; Anophelinae; Anopheles.

1 (bases 1 to 590)

Hong,Y.S., Hogan,J.R., Wang,X., Sarkar,A., Sim,C., Loftus,B.J.,



TITLE  
JOURNAL  
COMMENT

Fujisawa, T. and Gojobori, T.  
Hydra EST project  
Unpublished (2003)  
Contact: Jung Shan Hwang  
Center for Information Biology and DNA Data Bank of Japan  
National Institute of Genetics  
1111 Yata, Mishima, Shizuoka 411-8540, Japan  
Tel: 81-55-981-6847 (ex.6898)  
Fax: 81-55-981-6848  
Email: jhwang@lab.nig.ac.jp, URL: http://www.cib.nig.ac.jp.

FEATURES  
source

1..619  
Location/Qualifiers  
/organism="Hydra magnipapillata"  
/mol\_type="mRNA"  
/strain="105"  
/db\_xref="taxon:6085"  
/clone="hmp.17065"  
/tissue\_type="whole body"  
/dev\_stage="adult budding stage"  
/clone\_lib="Hydra magnipapillata cDNA library"

Query Match 17.8%; Score 34.2; DB 3; Length 619;  
Best Local Similarity 50.9%; Pred. No. 12; Mismatches 78; Indels 0; Gaps 0;  
Matches 81; Conservative 0;

ORIGIN

17 ATGAAGTGGCTTGACAAAGCATATCTTGTCTGATTCGTCTATTTCATACCTTCGGGGA 76  
|||||  
30 ATGAAGTGGCTTCAAGAGAACATCTATAGAAAGCAGATGTCAGAGTCTACAAA 89  
|||||  
77 ATAGATGTGAACCCCTTATAAACCGGGTTTCGCGAAGCATGCGCTAGTATCAT 136  
|||||  
90 ATAGAAGCAAGATATCCGGATCGAATTCCTGTGTAGTTGAAAAGCTCCTCGTCCACA 149  
|||||  
137 GATGACAAATGACGACTAAGCAAGAGTCTGTCCCTCGA 175  
|||||  
150 ATTCAGATATTGACAAAGAAAGTTCTGTGTCCCTCA 188  
|||||

RESULT 9  
DN244174 651 bp mRNA linear EST 29-JUN-2005  
LOCUS  
DEFINITION  
ACAB-aaa24nl5.g1 Hydra EST UCI 5 Hydra magnipapillata cDNA 5'  
similar to gb|AA045172.1| hypothetical GABA(A) receptor-associated  
protein [Branchiostoma belcheri], mRNA sequence.

ACCESSION  
DN244174  
VERSION  
DN244174.2 GI:68337862  
KEYWORDS  
EST.  
SOURCE  
Hydra magnipapillata  
Hydra magnipapillata  
Eukaryota; Metazoa; Cnidaria; Hydrozoa; Hydroidea; Anthomedusae;  
Hydridae; Hydra.

REFERENCE  
1 (bases 1 to 651)  
Bode,H., Blumberg,B., Steele,R., Wigge,P., Gee,L., Nguyen,Q.,  
Martinez,D., Kibler,D., Hampson,S., Clifton,S., Pape,D., Marx,M.,  
Hillier,L., Martin,J., Wylie,T., Dente,M., Theising,B., Bowers,Y.,  
Gibbons,M., Ritter,E., Bennett,J., Ronko,I., Tsagarishvili,R.,  
Maguire,L., Kennedy,S., Waterston,R. and Wilson,R.

WashU Hydra EST Project  
Unpublished (2002).  
On Mar 1, 2005 this sequence version replaced gi:60406628.  
Contact: Hans Bode  
WashU Hydra EST Project  
Washington University School of Medicine  
4444 Forest Park Parkway, Box 8501, St. Louis, MO 63108, USA  
Tel: 314 286 1800  
Fax: 314 286 1810

EMAIL: est@wustl.edu  
Library material provided by Hans Bode & Dirk Lindgens, Univ. of  
Calif., Irvine Library constructed by Dirk Lindgens, Univ. of  
Calif., Irvine Library sequenced by Washington University Genome  
Sequencing Center for information on obtaining a clone please  
contact: Hans Bode (hrobode@uci.edu)

This trace has been recalled with phred  
original value before phred recall for SL was 100  
original value before phred recall for SR was 790  
Seq primer: -40RP from Gibco.

FEATURES  
source

Location/Qualifiers  
1..651  
/organism="Hydra magnipapillata"  
/mol\_type="mRNA"  
/strain="105"  
/db\_xref="taxon:6085"  
/la5\_host="DH10B"  
/clone\_lib="Hydra EST UCI 5"  
/notes="vector: pSPORT1; Site\_1: Not I; Site\_2: Sal I;  
a.1st strand cDNA was primed with a Not I primer-adaptor  
(5' - PGACTAGTCTAGATCGAGCGGCCGCTT15-3')  
b.Double-stranded cDNA was ligated into the pSPORT 1-vector  
digested with Not I and cloned into the pSPORT 1-vector  
pre-cut with Not I and Sal I. c.The ligation mix was  
transformed into DH10B cells. d.The cells were grown in  
SOC = 5g yeast, 20g tryptone, 0.5 g NaCl, 10 mM MgSO4, 10  
mM MgCl, 0.2% glucose/liter, (no antibiotic). e.DMSO was  
added to a final conc. of 10% as a cryoprotectant and  
frozen f.The titre before freezing was determined as  
~2400/100 ul. Assuming a 10% loss upon freezing, the titre  
is probably ~2100/ 100 ul. g.9 tubes each containing ~  
2100 clones/100 ul (= total of ~19,000) are enclosed.  
h.The frequency of vectors containing inserts is 96% as  
determined by digestion check after picking 24 clones,  
miniprep and subsequent digestion with Not I and Sal I.  
i.A low level of 32P was used in the cDNA synthesis  
procedure. The level measured by holding a Geiger Counter  
next to a tube was background."

Query Match 17.8%; Score 34.2; DB 8; Length 651;  
Best Local Similarity 50.9%; Pred. No. 12; Mismatches 78; Indels 0; Gaps 0;  
Matches 81; Conservative 0;

ORIGIN

17 ATGAAGTGGCTTGACAAAGCATATCTTGTCTGATTCGTCTATTTCATACCTTCGGGGA 76  
|||||  
81 ATGAAGTGGCTTCAAGAGAACATATTTTGAAGAGCAGATGTCAGAGTCTACAAA 140  
|||||  
77 ATAGATGTGAACCCCTTATAAACCGGGTTTCGCGAAGCATGCGCTAGTATCAT 136  
|||||  
141 ATAGAAGCAAGATATCCGGATCGAATTCCTGTGTAGTTGAGAAAGCTCCTCGTCCACA 200  
|||||  
137 GATGACAAATGACGACTAAGCAAGAGTCTGTCCCTCGA 175  
|||||  
201 ATTCAGATATTGACAAAGAAAGTTTCTGTGTCCCTCA 239  
|||||

RESULT 10

CR281388

LOCUS

DEFINITION

CR281388 Oryza sativa library (Han B) Oryza sativa cDNA clone

Y620d12p5, mRNA sequence.

ACCESSION

CR281388

VERSION

CR281388.1 GI:44667954

KEYWORDS

EST.

SOURCE

Oryza sativa

Oryza sativa

Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;

SpERMatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;

Ehrhartoidae; Oryzaceae; Oryza.

REFERENCE

1 (bases 1 to 878)

AUTHORS

Han,B., Feng,Q., Huang,Y.C., Ying,K., Li,Y., Guan,J.P., Zhu,J.J.,

Zhao,Q., Hu,X., Liu,Y.L., Mu,J., Yu,Z., Chen,L., Fan,D.L.,

Weng,Q.J., Zhang,L., Lu,Y.Q., Yu,S.L., Liu,X.H., Lu,T.T.,

Zhang,Y.J., Lu,Y., Li,C., Li,T., Zhang,Y., Hu,H., Jia,P.X.,

Zhang,L., Lan,L.F., Chen,W., Wu,S.A. and Xue,Y.B.

Rice cDNA EST clone

Unpublished (2003)

JOURNAL

COMMENT

Contact: Han Bin

National Center for Gene Research  
Chinese Academy of Sciences  
500# Cao Bao Road, Shanghai 200233, China  
Email: bhan@ncgr.ac.cn  
Clone requests: bhan@ncgr.ac.cn  
This is rice cDNA est clone  
Web site: http://www.ncgr.ac.cn.

# FEATURES

Location/Qualifiers  
1. .878  
/organism="Oryza sativa"  
/mol\_type="mRNA"  
/db\_xref="taxon:4530"  
/clone="v620d12p5"  
/clone\_lib="Oryza sativa library (Han B)"

## ORIGIN

Query Match 17.8%; Score 34.2; DB 7; Length 878;  
Best Local Similarity 49.2%; Pred. No. 13;  
Matches 90; Conservative 0; Mismatches 93; Indels 0; Gaps 0;  
QY 2 CTGGGGCGGGCCATGAAGTGGCTTGACAGCATATCTTGTCTGATTGCTCTATTTT 61  
|||||  
Db 526 CTGGCTATTGTACCCATGAGGCTGACTATTGACGAGAAGTCGGGTTTACTGTGTATCAG 585  
QY 62 CAATACCTTCGGGGAATAGATGGAACCCCTTATATAACGCGGGTTTTCGAGAAACA 121  
|||||  
Db 586 GATTGGGCTTCGTGATAAAATCTTAAATTTGGGACAGCGGTATTTTCGTCATGG 645  
QY 122 TGGCGTAGTATCATTCATGACACATGAGCTAAGCAAAAGTGTGTCCTCGACCCCAAG 181  
|||||  
Db 646 CCCTCTGGTATAAAATCTGAGACACTTTACACGCTACAGAAGGTACCCCTCGACAGCA 705  
QY 182 AAG 184  
||  
Db 706 TAG 708

RESULT 11  
CNS079HE/c 945 bp DNA linear GSS 08-JUL-2001  
LOCUS T3 end of clone BB0AA029A02 of library BB0AA from strain CBS 4732  
DEFINITION of Pichia angusta, genomic survey sequence.

ACCESSION AL435288.1 GI:12218701

VERSION GSS.

KEYWORDS Pichia angusta

SOURCE Pichia angusta

ORGANISM Eukaryota; Fungi; Ascomycota; Saccharomycotina; Saccharomycetes;

Saccharomycetales; Saccharomycetaceae; Pichia.

REFERENCE 1 (bases 1 to 945)

AUTHORS Souciet,J.L., Aigle,M., Artiguenave,F., Blandin,G.,

Bolotin-Fukuhara,M., Bon,E., Brottier,P., Casaregola,S.,

de-Montigny,J., Dujon,B., Durrens,P., Lepingle,A., Llorente,B.,

Malpertuy,A., Neuveglise,C., Ozier-Kalogeropoulos,O., Potier,S.,

Saurin,W., Tekaiia,F., Toffano-Nioche,C., Wesolowski-Louvel,M.,

Wincker,P. and Weissenbach,J.

Genomic exploration of the hemiascomycetous yeasts: 1. A set of

yeast species for molecular evolution studies

FEBS Lett. 487 (1), 3-12 (2000)

JOURNAL 11152876

PUBMED 2 (bases 1 to 945)

REFERENCE Blandin,G., Llorente,B., Malpertuy,A., Wincker,P., Artiguenave,F.

AUTHORS and Dujon,B.

Genomic exploration of the hemiascomycetous yeasts: 13. Pichia

angusta

FEBS Lett. 487 (1), 76-81 (2000)

JOURNAL 11152888

PUBMED 3 (bases 1 to 945)

REFERENCE Genoscope.

AUTHORS Direct Submission

TITLE Submitted (08-SEP-2000) Genoscope - Centre National de Sequencage,

JOURNAL 2 rue Gaston Cremieux, CP 5706, 91057 EVRY cedex, FRANCE. (E-mail :

seqref@genoscope.cns.fr - Web : www.genoscope.cns.fr)

## COMMENT

This GSS is part of a random genomic sequencing program of thirteen yeast species: Saccharomyces bayanus var. uvarum, Saccharomyces exiguus, Saccharomyces servazzii, Zygosaccharomyces rouxii, Saccharomyces kluyveri, Kluyveromyces thermotolerans, Kluyveromyces lactis var. lactis, Kluyveromyces marxianus var. marxianus, Pichia angusta, Debaryomyces hansenii var. hansenii, Pichia sorbitophila, Candida tropicalis and Yarrowia lipolytica. Genomic inserts of 3 to 5 kb were prepared and both extremities were sequenced. See keywords for description of this sequence and for the sequence of the other extremity of this insert.

## FEATURES

Location/Qualifiers  
1. .945  
/organism="Pichia angusta"  
/mol\_type="genomic DNA"  
/strain="CBS 4732"  
/db\_xref="taxon:4905"  
/clone="BB0AA029A02"  
/clone\_lib="BB0AA"  
/notes="end : T3"  
misc\_feature <4. .>944  
/note="similar to Saccharomyces cerevisiae ORF YMR290c (HAS1 ; helicase associated with Set1p )  
1 putative frameshift(s)"  
/evidence=not\_experimental

## ORIGIN

Query Match 17.6%; Score 33.8; DB 11; Length 945;  
Best Local Similarity 56.3%; Pred. No. 18;  
Matches 62; Conservative 0; Mismatches 47; Indels 0; Gaps 0;  
QY 35 CATAACTCTGTCTGATTGCTCTATTTTCAATACCTTCGGGGAATAGATGTGAAACCCCT 94  
|||||  
Db 387 CAGAAATCTGTCTGCTTCGTCATATCAATGCTTCAAGTCTTAAAGATAAATCCCTT 328  
QY 95 TATATAACGCGGGTTCGAGAAACATCGCGCTAGTATCATTTGATGACA 143  
|||||  
Db 327 CGTATTTTGCAGTGATCCAGCAATCTACCTGGAGTGCAATGATCAAA 279

## RESULT 12

CNS0757U/c

LOCUS

DEFINITION

clone BA0AB033R09 of library BA0AB from strain CLIB 210 of

Kluyveromyces lactis, genomic survey sequence.

ACCESSION AL429760.1 GI:12212954

VERSION GSS.

KEYWORDS Kluyveromyces lactis

SOURCE Kluyveromyces lactis

ORGANISM Eukaryota; Fungi; Ascomycota; Saccharomycotina; Saccharomycetes;

Saccharomycetales; Saccharomycetaceae; Kluyveromyces.

REFERENCE 1 (bases 1 to 845)

AUTHORS Souciet,J.L., Aigle,M., Artiguenave,F., Blandin,G.,

Bolotin-Fukuhara,M., Bon,E., Brottier,P., Casaregola,S.,

de-Montigny,J., Dujon,B., Durrens,P., Lepingle,A., Llorente,B.,

Malpertuy,A., Neuveglise,C., Ozier-Kalogeropoulos,O., Potier,S.,

Saurin,W., Tekaiia,F., Toffano-Nioche,C., Wesolowski-Louvel,M.,

Wincker,P. and Weissenbach,J.

Genomic exploration of the hemiascomycetous yeasts: 1. A set of

yeast species for molecular evolution studies

FEBS Lett. 487 (1), 3-12 (2000)

JOURNAL 11152876

PUBMED 2 (bases 1 to 845)

REFERENCE Bolotin-Fukuhara,M., Toffano-Nioche,C., Artiguenave,F.,

AUTHORS Duchateau-Nguyen,G., Lemaire,M., Marmeis,R., Montrocher,R.,

Robert,C., Termier,M., Wincker,P. and Wesolowski-Louvel,M.

Genomic exploration of the hemiascomycetous yeasts: 11.

TITLE Kluyveromyces lactis

FEBS Lett. 487 (1), 66-70 (2000)

JOURNAL 11152886

PUBMED 3 (bases 1 to 845)

REFERENCE Genoscope.

AUTHORS Direct Submission

TITLE

Submitted (08-SEP-2000) Genoscope - Centre National de Sequencage, 2 rue Gaston Cremieux, CP 5706, 91057 EVRY cedex, FRANCE. (E-mail : segre@genoscope.cns.fr - Web : [www.genoscope.cns.fr](http://www.genoscope.cns.fr)) This GSS is part of a random genomic sequencing program of thirteen yeast species: *Saccharomyces bayanus* var. *uvarum*, *Saccharomyces exiguus*, *Saccharomyces servazzii*, *Zygosaccharomyces rouxii*, *Saccharomyces kluyveri*, *Kluyveromyces thermotolerans*, *Kluyveromyces lactis* var. *lactis*, *Kluyveromyces marxianus* var. *marxianus*, *Pichia angusta*, *Debaryomyces hansenii* var. *hansenii*, *Pichia sorbitophila*, *Candida tropicalis* and *Yarrowia lipolytica*. Genomic inserts of 3 to 5 kb were prepared and both extremities were sequenced. See keywords for description of this sequence and for the sequence of the other extremity of this insert.

Email: jbedell@iongenomics.com  
Plate: fabb001f081 row: a column: 17  
Seq primer: f Forward  
Class: methylation filtered  
High quality sequence stop: 463.  
Location/Qualifiers  
1..463  
/organism="Sorghum bicolor"  
/mol\_type="genomic DNA"  
/cultivar="ATx623"  
/db\_xref="taxon:4558"  
/clone="fabb001f081a17"  
/clone\_lib="Sorghum methylation filtered library (LibID:  
104)"  
/note="Organ: leaf; Vector: pBSK(-); Site 1: HincII; DNA  
prepared from purified nuclei was randomly sheared,  
end-repaired, size fractionated to enrich for the 0.5 to 5  
kb fraction, ligated into HincII-digested pBSK(-) vector  
and electroporated into E. coli cells. This is a  
methylation filtered library."

**ORIGIN**

Query Match	17.4%;	Score 33.4;	DB 10;	Length 463;
Best Local Similarity	57.0%;	Pred. No. 20;	Mismatches	46;
Matches	61;	Conservative	0;	Gaps 0;

```

Qy      62 CAATACCTTCGGGGAATAGATGTGAAAACCCCTTATAAACGGGGTTTTCGCAGAAACA 121
          ||| | | | | | | | | | | | | | | | | | | | | | | | | | | |
Db      354 CAGGCATTGTAGTAAGTGCTCGATAGCCTCCAAGGCCCTTGCCCTTTGCTGAAGC 295
          ||| | | | | | | | | | | | | | | | | | | | | | | | | | | |

Qy     122 TGGCGTAGTATCATGTATGACACATCGACTAAGCAAAAAGTGCTTGT 168
          ||| | | | | | | | | | | | | | | | | | | | | | | | | | | |
Db     294 GGCCATGGTTTCAAAGGTGACAGGTGAACCAGGCAAGCTGGGGTGT 248
          ||| | | | | | | | | | | | | | | | | | | | | | | | | | | |

```

**RESULT 14**  
**CA394392**  
**LOCUS**  
**DEFINITION**  
CA394392 494 bp mRNA linear EST 06-NOV-2002  
cs51a07.y1 Human Retinal pigment epithelium/choroid cDNA  
(Un-normalized, unambiguated): cs Homo sapiens cDNA clone cs51a07  
5', mRNA sequence.

**ACCESSION**  
VERSION  
KEYWORDS  
SOURCE  
ORGANISM

**REFERENCE**  
**AUTHORS**  
Wistow G., Bernstein S.L., Wyatt M.K., Farries R.N., Behal A.,  
Touchman J.W., Bouffard G., Smith D. and Peterson K.  
Expressed sequence tag analysis of human RPE/choroid for the  
NIH Bank Project: Over 600 non-redundant transcripts, novel genes  
and splice variants  
Mol. Vis. 8 (4), 205-220 (2002)  
12107410

**JOURNAL**  
**PUBLISHED**  
**COMMENT**

Contact: Wistow G  
Section on Molecular Structure and Function  
National Eye Institute  
6/331, NIH, Bethesda, MD 20892-2740, USA  
Tel: 301 402 3452  
Fax: 301 496 0078  
Email: graeme@helix.nih.gov  
Plate: 51 row: a column: 07  
Seq primer: MJRP1 reverse primer (AB1).  
Location/Qualifiers  
1..494  
/organism="Homo sapiens"  
/mol\_type="mRNA"  
/db\_xref="taxon:9606"  
/clone="cs51a07"  
/tissue type="RPE/choroid"

**FEATURES**  
source



```

/dev_stage="Adult"
/lab_host="EMDH10B"
/clone_lib="Human Retinal pigment epithelium/choroid cDNA
(Un-normalized, unamplified): cs"
/notes="Organ: Eye; Vector: pCMVSPORT6; Two different donor
eyes (75-80 years old) yielded approximately 600 mg of
dissected RPE/choroid tissue. This in turn yielded 340 ug
of total RNA and 7 ug of mRNA. A directionally cloned cDNA
library in the pCMVSPORT6 vector was constructed at Life
Technologies (Rockville, MD; now part of Invitrogen Corp),
essentially following the protocols of the SuperScript
Plasmid System (Invitrogen Corp).
<http://www.invitrogen.com/>). The library code
designation was cs. For this library, cDNA inserts were
cloned into the NotI/MluI sites of the vector. EST
analysis was performed on the unamplified library at the
NIH Intramural Sequencing Center (NISC)."
```

## ORIGIN

```

Query Match          17.4%; Score 33.4; DB 6; Length 494;
Best Local Similarity 52.5%; Pred. No. 21;
Matches 73; Conservative 0; Mismatches 66; Indels 0; Gaps 0;

QY 42 TTGCTGATTGCTGCTATTTTCAATACCTTCGGGGAATAGATGTGAAACCCCTTATAAAA 101
Db 7 TTTTCTACTTTGCTGCTCTCTACAGCTACAGCAATGAATCTGCTCAGCAGATGAAG 66

QY 102 CGCGGGTTTTTCGAGAAACATGCGCTAGTATCATTTGATGACAAATGAGCTAAGCAAAAAG 161
Db 67 CCTATTTTTCAGCATAAATATGCTTGCCATGATTAATAATAGAAATCAGCTAAGCATGAT 126

QY 162 TGCTTGTCCCTGACCCAA 180
Db 127 TTCTTTACTGGTGGACCAA 145
```

## RESULT 15

```

BH885133
LOCUS          728 bp      DNA      linear      GSS 05-AUG-2002
DEFINITION    hw61c11.b1 WGS-Zmay5F (JM107 adapted methyl filtered) Zea mays
               genomic clone hw61c11 5', genomic survey sequence.
ACCESSION     BH885133
VERSION       BH885133.1 GI:22121030
KEYWORDS      GSS.
SOURCE        Zea mays
ORGANISM      Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
               Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae; PACCAD
               clade; Panicoideae; Andropogoneae; Zea.
REFERENCE     1 (bases 1 to 728)
AUTHORS       Rabinowicz,P.D., O'Shaughnessy,A.L., Ballja,V., Dedhia,N.,
               Katzenburger,F., King,L., Miller,B., Muller,S., Nascimento,L.,
               Zutavern,T., McCombie,W.R. and Martienssen,R.A.
TITLE         Genomic shotgun sequences from Zea mays (methyl-filtered)
JOURNAL       Unpublished (2002)
COMMENT       Contact: W. Richard McCombie
               Lita Annenberg Hazen Genome Sequencing Center
               Cold Spring Harbor Laboratory
               PO Box 100, Cold Spring Harbor, NY 11724, USA
               Tel: 516 367 8884
               Fax: 516 367 8874
               Email: mcombie@cshl.org
               Plate: hw61 row: C column: 11
               Seq primer: -21M13UnivFwd
               Class: shotgun
               High quality sequence stop: 728.
```

## FEATURES source

```

1..728
/organism="Zea mays"
/mol_type="genomic DNA"
/cultivar="B73"
/db_xref="taxon:4577"
/clone="hw61c11"
```

```

/lab_host="JM107 or DH5a"
/clone_lib="WGS-Zmay5F (JM107 adapted methyl filtered)"
/notes="Organ: immature ears; Site_1: Xba I; Site_2: Xba I;
The vector was digested with XbaI and one nucleotide was
added by fill in in the recessive 3' end. The genomic DNA
was nebulized, end repaired, adaptor ligated and size
fractionated using sephadex. The resulting fragments were
between 0.8 and 3 kb and were cloned into the vector
(.x/y reads in M13mp19, b/g reads in pUC19). The same
ligation was transformed in either JM107 or DH5a."
```

## ORIGIN

```

Query Match          17.4%; Score 33.4; DB 9; Length 728;
Best Local Similarity 62.7%; Pred. No. 23;
Matches 52; Conservative 0; Mismatches 31; Indels 0; Gaps 0;

QY 110 TTGCGAGAAACATGCGCTAGTATCATTTGATGACAAACATGAGCTTAAGCAAAAGTGTGTC 169
Db 171 TTCATAGAGACATGCGATAATAGCAGGATACAAATAAGAAATAATTTCATCGTCTCTTC 230

QY 170 CCTGACCCCAAGAGGATGCTTT 192
Db 231 CTCCTCTTCAAGCAGGACCTAT 253
```

Search completed: March 7, 2006, 13:19:34  
Job time : 2366.02 secs



Result No.	Score	Query		Length	DB	ID	Description
		Match	%				
1	32.6	17.0	278	3	US-09-513-9990-31601		Sequence 31601, A
2	31.2	16.2	22404	3	US-09-949-016-11765		Sequence 11765, A
C 2	31.2	16.2	22404	3	US-09-949-016-15398		Sequence 15398, A
C 3	31.2	16.2	26167	3	US-09-454-466-1		Sequence 1, Appli
C 4	31.2	16.2	26167	3	US-09-949-016-13964		Sequence 13964, A
C 5	31.2	16.2	46745	3	US-09-949-016-48509		Sequence 48509, A
C 6	30.6	15.9	601	3	US-09-949-016-13175		Sequence 13175, A
C 7	30.6	15.9	58768	3	US-09-949-016-13175		Sequence 13175, A
C 8	30.4	15.8	1569	3	US-09-710-279-603		Sequence 603, App
C 9	30.4	15.8	3204	3	US-09-710-279-3485		Sequence 3485, Ap
C 10	29.8	15.5	10303	3	US-09-634-238-410		Sequence 410, App
C 11	29.8	15.5	99797	3	US-09-949-016-15255		Sequence 15255, A
C 12	29.6	15.4	601	3	US-09-949-016-187277		Sequence 187277, A
C 13	29.6	15.4	36855	3	US-09-949-016-17095		Sequence 17095, A
C 14	29.4	15.3	768	3	US-09-134-000C-3331		Sequence 3331, Ap
C 15	29.2	15.2	601	3	US-09-949-016-142351		Sequence 142351, A
C 16	29.2	15.2	2385	3	US-09-949-016-4210		Sequence 4010, Ap
C 17	29.2	15.2	3077	3	US-09-461-912A-20		Sequence 27, Appl
C 18	29.2	15.2	3092	2	US-08-426-627-3		Sequence 3, Appli
C 19	29.2	15.2	3126	2	US-08-477-396A-3		Sequence 3, Appli
C 20	29.2	15.2	3253	2	US-08-426-627-5		Sequence 5, Appli
C 21	29.2	15.2	4823	3	US-09-949-016-15769		Sequence 15769, A
C 22	29.2	15.2	160552	3	US-09-593-828-11		Sequence 11, Appl
C 23	29	15.1	601	3	US-09-949-016-44696		Sequence 44696, A
C 24	29	15.1	832	3	US-09-621-976-2813		Sequence 2813, Ap

; Patent No. 6812339  
; GENERAL INFORMATION:  
; APPLICANT: VENTER, J. Craig et al.  
; TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED  
; FILE REFERENCE: CL001307  
; CURRENT APPLICATION NUMBER: US/09/949,016  
; CURRENT FILING DATE: 2000-04-14  
; PRIOR APPLICATION NUMBER: 60/241,755  
; PRIOR FILING DATE: 2000-10-20  
; PRIOR APPLICATION NUMBER: 60/237,768  
; PRIOR FILING DATE: 2000-10-03  
; PRIOR APPLICATION NUMBER: 60/231,498  
; PRIOR FILING DATE: 2000-09-08  
; NUMBER OF SEQ ID NOS: 207012  
; SOFTWARE: FastSEQ for Windows Version 4.0  
; SEQ ID NO 11765  
; LENGTH: 22404  
; TYPE: DNA  
; ORGANISM: Human  
; FEATURE:  
; NAME/KEY: misc\_feature  
; LOCATION: (1)...(22404)  
; OTHER INFORMATION: n = A,T,C or G  
US-09-949-016-11765

Query Match 16.2%; Score 31.2; DB 3; Length 22404;  
Best Local Similarity 60.7%; Pred. No. 3.8;  
Matches 51; Conservative 0; Mismatches 33; Indels 0; Gaps 0;

QY 71 CGGGGAATAGATGTGAAAACCCCTTATAAAACGGGGTTTCGCAGAAACATCGCGTAGT 130

Db 19171 CGAGCAGAGAGATGAAAGGCGAGAGATGTGGAGGTGTCCAGACACAGCTGGGCGAGG 19112

QY 131 ATCATTGATGACAAACATGGACTAA 154  
Db 19111 AACACCCATGAGAACAGGGACTCA 19088

## RESULT 3

US-09-949-016-15398/c  
; Sequence 15398, Application US/09949016  
; Patent No. 6812339  
; GENERAL INFORMATION:  
; APPLICANT: VENTER, J. Craig et al.  
; TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED  
; FILE REFERENCE: CL001307  
; CURRENT APPLICATION NUMBER: US/09/949,016  
; CURRENT FILING DATE: 2000-04-14  
; PRIOR APPLICATION NUMBER: 60/241,755  
; PRIOR FILING DATE: 2000-10-20  
; PRIOR APPLICATION NUMBER: 60/237,768  
; PRIOR FILING DATE: 2000-10-03  
; PRIOR APPLICATION NUMBER: 60/231,498  
; PRIOR FILING DATE: 2000-09-08  
; NUMBER OF SEQ ID NOS: 207012  
; SOFTWARE: FastSEQ for Windows Version 4.0  
; SEQ ID NO 15398  
; LENGTH: 22404  
; TYPE: DNA  
; ORGANISM: Human  
; FEATURE:  
; NAME/KEY: misc\_feature  
; LOCATION: (1)...(22404)  
; OTHER INFORMATION: n = A,T,C or G  
US-09-949-016-15398

Query Match 16.2%; Score 31.2; DB 3; Length 22404;  
Best Local Similarity 60.7%; Pred. No. 3.8;  
Matches 51; Conservative 0; Mismatches 33; Indels 0; Gaps 0;

QY 71 CGGGGAATAGATGTGAAAACCCCTTATAAAACGGGGTTTCGCAGAAACATCGCGTAGT 130

Db 19171 CGAGCAGAGAGATGAAAGGCGAGAGATGTGGAGGTGTCCAGACACAGCTGGGCGAGG 19112  
QY 131 ATCATTGATGACAAACATGGACTAA 154  
Db 19111 AACACCCATGAGAACAGGGACTCA 19088

## RESULT 4

US-09-454-466-1/c  
; Sequence 1, Application US/09454466  
; Patent No. 6858425  
; GENERAL INFORMATION:  
; APPLICANT: Heus, Joris Jan  
; APPLICANT: Pharming Intellectual Property B.V.  
; TITLE OF INVENTION: HUMAN ACID ALPHA GLUCOSIDASE GENE AND BOVINE ALPHA-S1  
; TITLE OF INVENTION: CASEIN GENE SEQUENCES  
; FILE REFERENCE: 016994-013720US  
; CURRENT APPLICATION NUMBER: US/09/454,466  
; CURRENT FILING DATE: 1999-12-03  
; EARLIER APPLICATION NUMBER: 60/110,859  
; EARLIER FILING DATE: 1998-12-04  
; EARLIER APPLICATION NUMBER: 60/122,550  
; EARLIER FILING DATE: 1999-03-02  
; NUMBER OF SEQ ID NOS: 5  
; SOFTWARE: PatentIn Ver. 2.1  
; SEQ ID NO 1  
; LENGTH: 26167  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
US-09-454-466-1

Query Match 16.2%; Score 31.2; DB 3; Length 26167;  
Best Local Similarity 60.7%; Pred. No. 4.1;  
Matches 51; Conservative 0; Mismatches 33; Indels 0; Gaps 0;

QY 71 CGGGGAATAGATGTGAAAACCCCTTATAAAACGGGGTTTCGCAGAAACATCGCGTAGT 130

Db 17958 CGAGCAGAGAGATGAAAGGCGAGAGATGTGGAGGTGTCCAGACACAGCTGGGCGAGG 17899

QY 131 ATCATTGATGACAAACATGGACTAA 154

Db 17898 AACACCCATGAGAACAGGGACTCA 17875

## RESULT 5

US-09-949-016-13964/c  
; Sequence 13964, Application US/09949016  
; Patent No. 6812339  
; GENERAL INFORMATION:  
; APPLICANT: VENTER, J. Craig et al.  
; TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED  
; FILE REFERENCE: CL001307  
; CURRENT APPLICATION NUMBER: US/09/949,016  
; CURRENT FILING DATE: 2000-04-14  
; PRIOR APPLICATION NUMBER: 60/241,755  
; PRIOR FILING DATE: 2000-10-20  
; PRIOR APPLICATION NUMBER: 60/237,768  
; PRIOR FILING DATE: 2000-10-03  
; PRIOR APPLICATION NUMBER: 60/231,498  
; PRIOR FILING DATE: 2000-09-08  
; NUMBER OF SEQ ID NOS: 207012  
; SOFTWARE: FastSEQ for Windows Version 4.0  
; SEQ ID NO 13964  
; LENGTH: 46745  
; TYPE: DNA  
; ORGANISM: Human  
; FEATURE:  
; NAME/KEY: misc\_feature  
; LOCATION: (1)...(46745)  
; OTHER INFORMATION: n = A,T,C or G  
US-09-949-016-13964

Query Match 16.2%; Score 31.2; DB 3; Length 46745;  
Best Local Similarity 54.3%; Pred. No. 5.1;  
Matches 63; Conservative 0; Mismatches 53; Indels 0; Gaps 0;  
QY 46 CTGATTGGTATTTTCAATACCTTCGGGAAATAGATGTGAAACCCCTATATAAAGCG 105  
DB 40974 CTGCTCCACTGCTGCATGACCTTAGGTGGCAAAAGTCTTACCTTTCTACCTTAAG 40915  
QY 106 GGTTCGAGAAACATCGCTAGTATCATTTGATGACAAATGACATGACAAAG 161  
DB 40914 GATGGGCAATTCCTCTTTTATGATATCTTCAGAACAGCTGTCTTAAGTAAAG 40859  
RESULT 6  
US-09-949-016-48509  
; Sequence 48509, Application US/09949016  
; Patent No. 6812339  
; GENERAL INFORMATION:  
; APPLICANT: VENTER, J. Craig et al.  
; TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED  
; WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF  
; FILE REFERENCE: CL001307  
; CURRENT APPLICATION NUMBER: US/09/949,016  
; CURRENT FILING DATE: 2000-04-14  
; PRIOR APPLICATION NUMBER: 60/241,755  
; PRIOR FILING DATE: 2000-10-20  
; PRIOR APPLICATION NUMBER: 60/237,768  
; PRIOR FILING DATE: 2000-10-03  
; PRIOR APPLICATION NUMBER: 60/231,498  
; PRIOR FILING DATE: 2000-09-08  
; NUMBER OF SEQ ID NOS: 207012  
; SOFTWARE: FastSeq for Windows Version 4.0  
; SEQ ID NO 48509  
; LENGTH: 601  
; TYPE: DNA  
; ORGANISM: Human  
US-09-949-016-48509  
Query Match 15.9%; Score 30.6; DB 3; Length 601;  
Best Local Similarity 58.1%; Pred. No. 1.5;  
Matches 54; Conservative 0; Mismatches 39; Indels 0; Gaps 0;  
QY 56 TATTTTCAATACCTTCGGGAAATAGATGTGAAACCCCTTATAAAGCGGGTTTCGCA 115  
DB 149 TACTTTCATTAACCTTTAGGAAGCTGAAGGGGAAATAATCAATAAGCAGCTCTCTTGACT 208  
QY 116 GAAACATGCGCTAGTATCATTTGATGACAAATG 148  
DB 209 CAAAGATGAGCTGAAATAATACATGAATGAATG 241  
RESULT 7  
US-09-949-016-13175/c  
; Sequence 13175, Application US/09949016  
; Patent No. 6812339  
; GENERAL INFORMATION:  
; APPLICANT: VENTER, J. Craig et al.  
; TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED  
; WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF  
; FILE REFERENCE: CL001307  
; CURRENT APPLICATION NUMBER: US/09/949,016  
; CURRENT FILING DATE: 2000-04-14  
; PRIOR APPLICATION NUMBER: 60/241,755  
; PRIOR FILING DATE: 2000-10-20  
; PRIOR APPLICATION NUMBER: 60/237,768  
; PRIOR FILING DATE: 2000-10-03  
; PRIOR APPLICATION NUMBER: 60/231,498  
; PRIOR FILING DATE: 2000-09-08  
; NUMBER OF SEQ ID NOS: 207012  
; SOFTWARE: FastSeq for Windows Version 4.0  
; SEQ ID NO 13175  
; LENGTH: 58768

TYPE: DNA  
ORGANISM: Human  
FEATURE:  
NAME/KEY: misc feature  
LOCATION: (1)... (58768)  
OTHER INFORMATION: n = A,T,C or G  
US-09-949-016-13175  
Query Match 15.9%; Score 30.6; DB 3; Length 58768;  
Best Local Similarity 58.1%; Pred. No. 9.1;  
Matches 54; Conservative 0; Mismatches 39; Indels 0; Gaps 0;  
QY 56 TATTTTCAATACCTTCGGGAAATAGATGTGAAACCCCTTATAAAGCGGGTTTCGCA 115  
DB 43027 TACTTTCATTAACCTTTAGGAAGCTGAAGGGGAAATAATCAATAAGCAGCTCTCTTGACT 42968  
QY 116 GAAACATGCGCTAGTATCATTTGATGACAAATG 148  
DB 42967 CAAAGATGAGCTGAAATAATACATGAATGAATG 42935  
RESULT 8  
US-09-710-279-603  
; Sequence 603, Application US/09710279  
; Patent No. 6703492  
; GENERAL INFORMATION:  
; APPLICANT: KIMMERLY, WILLIAM JOHN  
; TITLE OF INVENTION: STAPHYLOCOCCUS EPIDERMIDIS NUCLEIC ACIDS AND PROTEINS  
; FILE REFERENCE: PU3480US  
; CURRENT APPLICATION NUMBER: US/09/710,279  
; CURRENT FILING DATE: 2000-11-09  
; PRIOR APPLICATION NUMBER: 60/164,258  
; PRIOR FILING DATE: 1999-11-09  
; NUMBER OF SEQ ID NOS: 4472  
; SOFTWARE: PatentIn Ver. 2.1  
; SEQ ID NO 603  
; LENGTH: 1569  
; TYPE: DNA  
; ORGANISM: Artificial Sequence  
; FEATURE:  
; OTHER INFORMATION: Description of Artificial Sequence: synthetic  
; OTHER INFORMATION: nucleic acid sequence  
US-09-710-279-603  
Query Match 15.8%; Score 30.4; DB 3; Length 1569;  
Best Local Similarity 57.3%; Pred. No. 2.5;  
Matches 55; Conservative 0; Mismatches 41; Indels 0; Gaps 0;  
QY 49 ATTGCTTATTTTCAATACCTTCGGGAAATAGATGTGAAACCCCTTATAAAGCGGGT 108  
DB 87 ATATGAATATGAACAAATAATGAGCGATCCATATCTCTCAGCTTATAAAGCTAATGT 146  
QY 109 TTTCGAGAAACATGCGCTAGTATCATTTGATGACAA 144  
DB 147 TAACGCGGATATATTGTGATAGTTGATTGAATGA 182  
RESULT 9  
US-09-710-279-3485  
; Sequence 3485, Application US/09710279  
; Patent No. 6703492  
; GENERAL INFORMATION:  
; APPLICANT: KIMMERLY, WILLIAM JOHN  
; TITLE OF INVENTION: STAPHYLOCOCCUS EPIDERMIDIS NUCLEIC ACIDS AND PROTEINS  
; FILE REFERENCE: PU3480US  
; CURRENT APPLICATION NUMBER: US/09/710,279  
; CURRENT FILING DATE: 2000-11-09  
; PRIOR APPLICATION NUMBER: 60/164,258  
; PRIOR FILING DATE: 1999-11-09  
; NUMBER OF SEQ ID NOS: 4472  
; SOFTWARE: PatentIn Ver. 2.1  
; SEQ ID NO 3485  
; LENGTH: 3204

```
; TYPE: DNA
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: synthetic
US-09-710-279-3485

Query Match          15.8%; Score 30.4; DB 3; Length 3204;
Best Local Similarity 57.3%; Pred. No. 3.4; Indels 0; Gaps 0;
Matches 55; Conservative 0; Mismatches 41;

QY 49 ATTGCTCTATTTCCTCAATACCTTCGGGAAATAGATGTGAAACCCCTTATAAAACGGGGT 108
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 542 ATATGATATGACAAATAATATGAGCATCATCTCTCAGTCTATATAAACTAATGT 601

QY 109 TTTCGGCAGAAACATGCGCTAGTATCATATGATGACAA 144
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 602 TAACGGGGATATATTGATAGTTTGATTAAATGAAAA 637

RESULT 10
US-09-634-238-410/c
; Sequence 410, Application US/09634238
; Patent No. 6544772
; GENERAL INFORMATION:
; APPLICANT: Glenn, Matthew
; APPLICANT: Havukkala, Ilkka J.
; APPLICANT: Blokeberg, Leonard, N.
; APPLICANT: Lubbers, Mark W.
; APPLICANT: Dekker, James
; APPLICANT: Christenson, Anna C.
; APPLICANT: Holland, Ross
; APPLICANT: O'Toole, Paul W.
; APPLICANT: Reid, Julian R.
; APPLICANT: Coolbear, Timothy
; TITLE OF INVENTION: Polynucleotides, materials incorporating
; them and methods for using them.
; FILE REFERENCE: 11000.1043U1
; CURRENT FILING DATE: 2000-08-08
; NUMBER OF SEQ ID NOS: 422
; SOFTWARE: FastSeq for Windows Version 4.0
; SEQ ID NO 410
; LENGTH: 10303
; TYPE: DNA
; ORGANISM: Lactobacillus rhamnosus
US-09-634-238-410

Query Match          15.5%; Score 29.8; DB 3; Length 10303;
Best Local Similarity 60.5%; Pred. No. 8.6; Indels 0; Gaps 0;
Matches 49; Conservative 0; Mismatches 32;

QY 112 CGCAGAAACATGCGCTAGTATCATATGATGACAAACATGACTAAGCAAAAGTCTTGCCC 171
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 8088 CGAAGAAAGCAAGCAAGCATGTTGTAAGAAGTCATGACGAGCAAAAGAAATGTGCG 8029

QY 172 CTGACCCCAAGAGATGCTTT 192
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 8028 CGGCTCGAAAATGCGGCTTT 8008

RESULT 11
US-09-949-016-15255/c
; Sequence 15255, Application US/09949016
; Patent No. 6812339
; GENERAL INFORMATION:
; APPLICANT: VENTER, J. Craig et al.
; TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED
; WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF
; FILE REFERENCE: CL001307
; CURRENT FILING DATE: 2000-04-14
; PRIOR APPLICATION NUMBER: US/09/949,016
; PRIOR FILING DATE: 2000-04-14
```

```
; PRIOR FILING DATE: 2000-10-20
; PRIOR APPLICATION NUMBER: 60/237,768
; PRIOR FILING DATE: 2000-10-03
; PRIOR APPLICATION NUMBER: 60/231,498
; PRIOR FILING DATE: 2000-09-08
; NUMBER OF SEQ ID NOS: 207012
; SOFTWARE: FastSeq for Windows Version 4.0
; SEQ ID NO 15255
; LENGTH: 99797
; TYPE: DNA
; ORGANISM: Human
; NAME/KEY: misc feature
; LOCATION: (1)- (99797)
; OTHER INFORMATION: n = A,T,C or G
US-09-949-016-15255

Query Match          15.5%; Score 29.8; DB 3; Length 99797;
Best Local Similarity 51.1%; Pred. No. 21; Indels 0; Gaps 0;
Matches 70; Conservative 0; Mismatches 67;

QY 40 TCTTGCTGATTCGTCCTATTTCATATACCTTCGGGAAATAGATGTGAAACCCCTTATAA 99
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 32501 TGTAGGATGAATCCTGCTATTTCATGAGACTAGTGTTTAAAGGATGTGCAGGGGGTTAGAC 32442

QY 100 AACGCGGGTTTTCGCAGAAACATGCGCTAGTATCATATGATGACAAACATGACTAAGCAAA 159
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 32441 GGGCGGGGAGAGCAGTACAGGACAGTTACCAAGCCATAAAACACGAGCTGAAC 32382

QY 160 AGTGTCTGTCCCTGAC 176
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 32381 AATTCCTGCGCTCTGAC 32365

RESULT 12
US-09-949-016-187277/c
; Sequence 187277, Application US/09949016
; Patent No. 6812339
; GENERAL INFORMATION:
; APPLICANT: VENTER, J. Craig et al.
; TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED
; WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF
; FILE REFERENCE: CL001307
; CURRENT APPLICATION NUMBER: US/09/949,016
; CURRENT FILING DATE: 2000-04-14
; PRIOR APPLICATION NUMBER: 60/241,755
; PRIOR FILING DATE: 2000-10-20
; PRIOR APPLICATION NUMBER: 60/237,768
; PRIOR FILING DATE: 2000-10-03
; PRIOR APPLICATION NUMBER: 60/231,498
; PRIOR FILING DATE: 2000-09-08
; NUMBER OF SEQ ID NOS: 207012
; SOFTWARE: FastSeq for Windows Version 4.0
; SEQ ID NO 187277
; LENGTH: 601
; TYPE: DNA
; ORGANISM: Human
US-09-949-016-187277

Query Match          15.4%; Score 29.6; DB 3; Length 601;
Best Local Similarity 59.5%; Pred. No. 3.3; Indels 0; Gaps 0;
Matches 50; Conservative 0; Mismatches 34;

QY 96 ATAAACGGGGTTTTCGCAGAAACATGCGCTAGTATCATATGATGACAAACATGACTAAG 155
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 108 ATATGACCCAGTAGTCTTTGAAAGTCCTTCCTATCTTTTATGACAGATGTTCCAG 49

QY 156 CAAAGTGTGTCCTGACCA 179
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 48 CTTATTGTATATTCTGCCCCA 25

RESULT 13
```



***this Page Blank (uspto)***

GenCore version 5.1.7  
Copyright (c) 1993 - 2006 Bioceleration Ltd.

OM nucleic - nucleic search, using sw model

Run on: March 7, 2006, 12:14:00 ; Search time 526.953 Seconds  
(without alignments)  
3013.020 Million cell updates/sec

Title: US-10-782-899-1\_COPY\_1\_192

Perfect score: 192  
Sequence: 1 gctggcgccggcgccatga.....tgacccaagaagatgcttt 192

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 9793542 seqs, 4134689005 residues

Total number of hits satisfying chosen parameters: 19587084

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published Applications NA\_Main:\*  
1: /cgn2\_6/prodata/1/pubpna/us07\_PUBCOMB.seq:\*  
2: /cgn2\_6/prodata/1/pubpna/us08\_PUBCOMB.seq:\*  
3: /cgn2\_6/prodata/1/pubpna/us09A\_PUBCOMB.seq:\*  
4: /cgn2\_6/prodata/1/pubpna/us09B\_PUBCOMB.seq:\*  
5: /cgn2\_6/prodata/1/pubpna/us10A\_PUBCOMB.seq:\*  
6: /cgn2\_6/prodata/1/pubpna/us10B\_PUBCOMB.seq:\*  
7: /cgn2\_6/prodata/1/pubpna/us10C\_PUBCOMB.seq:\*  
8: /cgn2\_6/prodata/1/pubpna/us10D\_PUBCOMB.seq:\*  
9: /cgn2\_6/prodata/1/pubpna/us10E\_PUBCOMB.seq:\*  
10: /cgn2\_6/prodata/1/pubpna/us11\_PUBCOMB.seq:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	192	100.0	600	3	US-09-816-391A-1
2	192	100.0	600	8	US-10-782-899-1
3	175.2	91.2	2256646	7	US-10-470-565-1
4	34.6	18.0	425	8	US-10-425-115-60228
5	33.2	17.3	824	5	US-10-027-632-167810
6	33.2	17.3	824	6	US-10-027-632-167810
7	32.4	16.9	484	5	US-10-027-632-6585
8	32.4	16.9	484	6	US-10-027-632-6585
9	32.4	16.9	2717	10	US-11-097-143-30178
10	32	16.7	495269	7	US-10-398-221-8
11	32	16.7	3011208	7	US-10-398-221-2058
12	31.6	16.5	732	8	US-10-425-115-71957
13	31.6	16.5	1437	4	US-09-925-065A-691107
14	31.6	16.5	1437	4	US-09-925-065A-691108
15	31.2	16.2	360	8	US-10-674-124-12097
16	31.2	16.2	1485	4	US-09-925-065A-49312
17	31.2	16.2	1485	4	US-09-925-065A-49313
18	31.2	16.2	18272	3	US-09-764-891-8869
19	31	16.1	1771	4	US-09-925-065A-65588
20	30.8	16.0	639	4	US-09-925-065A-170277
21	30.8	16.0	639	4	US-09-925-065A-170278
22	30.6	15.9	466	7	US-10-021-323-5754
23	30.6	15.9	549	7	US-10-260-238-5041

ALIGNMENTS

RESULT 1

US-09-816-391A-1  
; Sequence 1, Application US/09816391A  
; Patent No. US2002005486SA1  
; GENERAL INFORMATION:  
; APPLICANT: FUJIMORI, Minoru  
; APPLICANT: TANIGUCHI, Shunichiro  
; APPLICANT: AMANO, Jun  
; APPLICANT: YAZAWA, Kazuyuki  
; APPLICANT: KANO, Yasunobu  
; APPLICANT: NAKAMURA, Toshiyuki  
; APPLICANT: SASAKI, Takayuki  
; TITLE OF INVENTION: Anaerobic bacterium as a drug for cancer gene therapy  
; FILE REFERENCE: 2001-WWC/01736  
; CURRENT APPLICATION NUMBER: US/09/816,391A  
; CURRENT FILING DATE: 2001-03-26  
; PRIOR APPLICATION NUMBER: JP 00/287688  
; PRIOR FILING DATE: 2000-09-21  
; NUMBER OF SEQ ID NOS: 3  
; SEQ ID NO 1  
; LENGTH: 600  
; TYPE: DNA  
; ORGANISM: Bifidobacterium longum  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (193)...(471)  
US-09-816-391A-1

Query Match 100.0%; Score 192; DB 3; Length 600;  
Best Local Similarity 100.0%; Pred. No. 4.7e-54;  
Matches 192; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 GCTGGCGCGCGGCCCATGAAGTGGCTTGCAAGCATATCTTGTCGATTGCTCTATTT 60  
DB 1 GCTGGCGCGCGGCCCATGAAGTGGCTTGCAAGCATATCTTGTCGATTGCTCTATTT 60  
QY 61 TCAATACCTTCGGGGAAATAGATGTGAAACCCCTTATAAAACGCGGGTTTTCGACAGAAC 120  
DB 61 TCAATACCTTCGGGGAAATAGATGTGAAACCCCTTATAAAACGCGGGTTTTCGACAGAAC 120  
QY 121 ATGCGCTAGTATCATTTGATGACAACTGACATGGAACAAAGTCTTGTCCCTGACCCAA 180  
DB 121 ATGCGCTAGTATCATTTGATGACAACTGACATGGAACAAAGTCTTGTCCCTGACCCAA 180  
QY 181 GAAGGATGCTTT 192  
|||||

Sequence 308556,  
Sequence 308557,  
Sequence 396377,  
Sequence 599874,  
Sequence 905740,  
Sequence 912520,  
Sequence 10536, A  
Sequence 10536, A  
Sequence 51874, A  
Sequence 16206, A  
Sequence 460994,  
Sequence 250, App  
Sequence 41046, A  
Sequence 185956,  
Sequence 185956,  
Sequence 684507,  
Sequence 684508,  
Sequence 25530, A  
Sequence 436, App  
Sequence 1207, Ap  
Sequence 6890, Ap

```
Db 181 GAAGGATGCTTT 192

RESULT 2
US-10-782-899-1
; Sequence 1, Application US/10782899
; Publication No. US20050025745A1
; GENERAL INFORMATION:
; APPLICANT: FUJIMORI, MINORU
; APPLICANT: TANIGUCHI, SHUNICHIRO
; APPLICANT: AMANO, JUN
; APPLICANT: YAZAWA, KAZUYUKI
; APPLICANT: KANO, YASUNOBU
; APPLICANT: NAKAMURA, TOSHIYUKI
; APPLICANT: SASAKI, TAKAYUKI
; TITLE OF INVENTION: ANAEROBIC BACTERIUM AS A DRUG FOR CANCER GENE THERAPY
; FILE REFERENCE: 671308-2003.1
; CURRENT APPLICATION NUMBER: US/10/782,899
; CURRENT FILING DATE: 2004-02-23
; PRIOR APPLICATION NUMBER: 09/816,391
; PRIOR FILING DATE: 2001-03-26
; PRIOR APPLICATION NUMBER: JP 2000-287688
; PRIOR FILING DATE: 2000-09-21
; NUMBER OF SEQ ID NOS: 4
; SOFTWARE: PatentIn Ver. 3.2
; SEQ ID NO 1
; LENGTH: 600
; TYPE: DNA
; ORGANISM: Bifidobacterium longum
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (193)..(471)
US-10-782-899-1

Query Match 100.0%; Score 192; DB 8; Length 600;
Best Local Similarity 100.0%; Pred. No. 4.7e-54;
Matches 192; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 GCTGGGGGGGGCCCATGAAGTGGCTTGACAAGCATATCTTGTCTGATTCGCTATTT 60
DB 1 GCTGGGGGGGGGGCCCATGAAGTGGCTTGACAAGCATATCTTGTCTGATTCGCTATTT 60

QY 61 TCAATACCTTCGGGGAATAGATGTGAAAACCCCTTATAAAACGGGGTTTTCGAGAAAC 120
DB 61 TCAATACCTTCGGGGAATAGATGTGAAAACCCCTTATAAAACGGGGTTTTCGAGAAAC 120

QY 121 ATCGGCTAGTATCATTTGATGACAAACATGGACTAAGCAAAAGTGTGTCCTGACCCAA 180
DB 121 ATCGGCTAGTATCATTTGATGACAAACATGGACTAAGCAAAAGTGTGTCCTGACCCAA 180

QY 181 GAAGGATGCTTT 192
DB 181 GAAGGATGCTTT 192

RESULT 3
US-10-470-565-1/c
; Sequence 1, Application US/10470565
; Publication No. US20040126870A1
; GENERAL INFORMATION:
; APPLICANT: Societe des Produits Nestle S.A.
; TITLE OF INVENTION: NCC2705 - the genome of a Bifidobacterium
; FILE REFERENCE: 80290/WO
; CURRENT APPLICATION NUMBER: US/10/470,565
; CURRENT FILING DATE: 2003-07-29
; PRIOR APPLICATION NUMBER: EP 01102050.0
; PRIOR FILING DATE: 2001-01-30
; NUMBER OF SEQ ID NOS: 2
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 1
; LENGTH: 2256646
; TYPE: DNA
; ORGANISM: Bifidobacterium longum

US-10-470-565-1
Query Match 91.2%; Score 175.2; DB 7; Length 2256646;
Best Local Similarity 97.4%; Pred. No. 1.2e-46;
Matches 189; Conservative 0; Mismatches 3; Indels 2; Gaps 1;

QY 1 GCTGGGGGGGGCCCATGAAGTGGCTTGACAAGCATATCTTGTCTGATTCGCTATTT 60
DB 2242172 GCTGGGGGGGGCCCATGAAGTGGCTTGACAAGCATATCTTGTCTGATTCGCTATTT 2242113

QY 61 TC--AATACCTTCGGGGAATAGATGTGAAAACCCCTTATAAAACGGGGTTTTCGAGAA 118
DB 2242112 TCATACCCCTTCGGGGAATAGATGTGAAAACCCCTTATAAAACGGGGTTTTCGAGAA 2242053

QY 119 ACATGCGCTAGTATCATTTGATGACACATGGACTAAGCAAAAGTGTGTCCTCGACCC 178
DB 2242052 ACATGCGCTAGTATCATTTGATGACACATGGACTAAGCAAAAGTGTGTCCTCGACCC 2241993

QY 179 AAGAAGGATGCTTT 192
DB 2241992 AAGAAGGATGCTTT 2241979

RESULT 4
US-10-425-115-60228/c
; Sequence 60228, Application US/10425115
; Publication No. US20040214272A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa, Thomas J.
; APPLICANT: Kovalic, David K.
; APPLICANT: Zhou, Yihua
; APPLICANT: Cao, Yongwei
; TITLE OF INVENTION: Nucleic Acid Molecules and Other Molecules Associated With
; TITLE OF INVENTION: Plants
; FILE REFERENCE: 38-21(S3222)B
; CURRENT APPLICATION NUMBER: US/10/425,115
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 369326
; SEQ ID NO 60228
; LENGTH: 425
; TYPE: DNA
; ORGANISM: Zea mays
; FEATURE:
; OTHER INFORMATION: Clone ID: MRT4577_15492C.1
US-10-425-115-60228

Query Match 18.0%; Score 34.6; DB 8; Length 425;
Best Local Similarity 51.6%; Pred. No. 0.6;
Matches 79; Conservative 0; Mismatches 74; Indels 0; Gaps 0;

QY 16 CATGAAGTGGCTTGACAAGCATATCTTGTCTGATTCGCTATTTTCAATACCTTCGGGG 75
DB 187 CTTGAAATCTACTGACAAAGATAAGACTAAATTTTACCCCAAAATACTATATATGC 128

QY 76 AAATAGATGTGAAAACCCCTTATAAAACGGGGTTTTCGAGAAACATGCGCTAGTATCAT 135
DB 127 AATGACAGCAATTAATACATTTGAATGCTATTTTACATATATACAAAGCACTGCTAGACT 68

QY 136 TGATGACACATGGACTAAGCAAAAGTGTGT 168
DB 67 TACTGCCAACAAATAGCTAAACAGAGAGGTGTT 35

RESULT 5
US-10-027-632-167810/c
; Sequence 167810, Application US/10027632
; Publication No. US20020198371A1
; GENERAL INFORMATION:
; APPLICANT: Wang, David G.
; TITLE OF INVENTION: Identification and Mapping of Single Nucleotide
; TITLE OF INVENTION: Polymorphisms in the Human Genome
; FILE REFERENCE: 108827.129
; CURRENT APPLICATION NUMBER: US/10/027,632
```



```

Qy 76 AAATAGATGTGAAAAACCCCTTATATAAACCGCGGGTTCCTCCAGAAACATCGCGCTAGTATCAT 130
Db 384 ATATATAATAGAAAAACCCAAAAGAGTGAAGCAATATTTTAAAGAAAGGTTGAACATGGATCTA 325

Qy 136 TGATGACAAACATGGACTAAGCAAAAGTCTTGTC 169
Db 324 TGATGAGAACAAAGTCATTAAGCAAGGGTTGTAATC 291

RESULT 7
US-10-027-632-6585
; Sequence 6585, Application US/10027632
; Publication No. US20020198371A1
; GENERAL INFORMATION:
; APPLICANT: Wang, David G.
; TITLE OF INVENTION: Identification and Mapping of Single Nucleotide
; POLYMORPHISMS IN THE HUMAN GENOME
; FILE REFERENCE: 108827.129
; CURRENT APPLICATION NUMBER: US/10/027,632
; CURRENT FILING DATE: 2002-04-30
; PRIOR APPLICATION NUMBER: US 60/218,006
; PRIOR FILING DATE: 2000-07-12
; PRIOR APPLICATION NUMBER: US 60/198,676
; PRIOR FILING DATE: 2000-04-20
; PRIOR APPLICATION NUMBER: US 60/193,483
; PRIOR FILING DATE: 2000-03-29
; PRIOR APPLICATION NUMBER: US 60/185,218
; PRIOR FILING DATE: 2000-02-24
; PRIOR APPLICATION NUMBER: US 60/167,363
; PRIOR FILING DATE: 1999-11-23
; PRIOR APPLICATION NUMBER: US 60/156,358
; PRIOR FILING DATE: 1999-09-28
; PRIOR APPLICATION NUMBER: US 60/146,002
; PRIOR FILING DATE: 1999-08-09

```

```

; SEQ ID NO 6585
; LENGTH: 484
; TYPE: DNA
; ORGANISM: Human
US-10-027-632-6585

Query Match      16.9%; Score 32.4; DB 5; Length 484;
Best Local Similarity 60.7%; Pred. NO. 3.6;
Matches 51; Conservative 1; Mismatches 32; Indels 0; Gaps 0

Qy      36  ATATCTTGTCGTGATTCGTCTATTTTCAATACCTTCGGGGAATAGATCTGAAAACCCCTT 95
Db      130 AGAATATCTCTTTTTCACACCTACTGCTTAATCCCTCCGGGCTAANTAGGCATGACAATAATA 189

Qy      96  ATAAAACGCGGGTTTTCGAGAA 119
Db      190 ATGAATACTTTTTTTCGAGAA 213

RESULT 8
US-10-027-632-6585
; Sequence 6585, Application US/10027632
; Publication No. US20030204075A9
; GENERAL INFORMATION:
; APPLICANT: Wang, David G.
; TITLE OF INVENTION: Identification and Mapping of Single Nucleotide
; TITLE OF INVENTION: Polymorphisms in the Human Genome
; FILE REFERENCE: 108827.129
; CURRENT APPLICATION NUMBER: US/10/027,632
; CURRENT FILING DATE: 2002-04-30
; PRIOR APPLICATION NUMBER: US 60/218,006
; PRIOR FILING DATE: 2000-07-12
; PRIOR APPLICATION NUMBER: US 60/198,676
; PRIOR FILING DATE: 2000-04-20
; PRIOR APPLICATION NUMBER: US 60/193,483
; PRIOR FILING DATE: 2000-03-29
; PRIOR APPLICATION NUMBER: US 60/185,218

```

Query Match	16.9%	Score 32.4;	DB 10;	Length 2717;
Best Local Similarity	56.6%;	Pred. No. 8.2;		
Matches 60;	Conservative	0;	Mismatches 46;	Indels 0;
Gaps	0			
Qy	39	ATCTTGTCGATTCTGCTATTTTTC	AATCCTTCGGGGAATAGATG	TAAGCCCTTATA 98
Db	369	ATAGTTTACAGATACATATAAAAT	TACAAATACATACAGCAAAAGCT	CATATATAGAGACTTAA 310

RESULT 11  
 US-10-398-221-2058  
 ; Sequence 2058, Application US/10398221  
 ; Publication No. US20040018514A1  
 ; GENERAL INFORMATION:  
 ; APPLICANT: KUNST, Frederik  
 ; APPLICANT: GLASER, Philippe  
 ; TITLE OF INVENTION: Listeria innocua, genome and applications  
 ; FILE REFERENCE: 344 702 - US  
 ; CURRENT APPLICATION NUMBER: US/10/398,221  
 ; CURRENT FILING DATE: 2003-03-27  
 ; PRIOR APPLICATION NUMBER: PCT/FR 01/03 061  
 ; PRIOR FILING DATE: 2001-10-04  
 ; PRIOR APPLICATION NUMBER: FR 00/12 697  
 ; PRIOR FILING DATE: 2000-10-04  
 ; NUMBER OF SEQ ID NOS: 4025  
 ; SOFTWARE: PatentIn version 3.0  
 ; SEQ ID NO 2058  
 ; LENGTH: 3011208  
 ; TYPE: DNA  
 ; ORGANISM: Listeria innocua  
 US-10-398-221-2058  
 Query Match 16.7%; Score 32; DB 7; Length 3011208;

```
Best Local Similarity 50.7%; Pred. No. 3.1e+02;
Matches 77; Conservative 0; Mismatches 75; Indels 0; Gaps 0;

QY 35 CATAATCTTGTCTGATTGCTTATTTTCAATACCTTCGGGGAATAGATGTGAAACCCCT 94
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 1529778 CATGAACCTGGACACGCTTCATCTTCATGTTATTCGGAGAGCCATTTGAAACACT 1529837

QY 95 TATAAACGCGGTTTTTCGCAGAAACATGCGGTAGTATCATTTGATGACCAACATGGACTAA 154
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 1529838 GATTAGCAATGAATGTTGCGGAACTGCTTCTACTTTTGCCTGAAATGATTATTCGCGAT 1529897

QY 155 GCAAAAGTGTCTTCCTCCCTGACCAAGAGGA 186
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 1529898 GCTTCTGTAAGAAGATGCCAAACAAAGAAGA 1529929

RESULT 12
US-10-425-115-71957/c
; Sequence 71957, Application US/10425115
; Publication No. US20040214272A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa, Thomas J.
; APPLICANT: Kovalic, David K.
; APPLICANT: Zhou, Yihua
; APPLICANT: Cao, Yongwei
; TITLE OF INVENTION: Nucleic Acid Molecules and Other Molecules Associated with
; FILE OF INVENTION: Plants
; FILE REFERENCE: 38-21(53222)B
; CURRENT APPLICATION NUMBER: US/10/425,115
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 369326
; SEQ ID NO 71957
; LENGTH: 732
; TYPE: DNA
; ORGANISM: Zea mays
; FEATURE:
; NAME/KEY: unsure
; LOCATION: (1)..(732)
; OTHER INFORMATION: unsure at all n locations
;
; OTHER INFORMATION: Clone ID: MRT4577_165627C.1
US-10-425-115-71957

Query Match 16.5%; Score 31.6; DB 8; Length 732;
Best Local Similarity 58.5%; Pred. No. 8.1;
Matches 55; Conservative 0; Mismatches 39; Indels 0; Gaps 0;

QY 54 TCTATTTTCATACCTTCGGGGAATAGATGTGAAACCCCTTATAAACCGCGGTTTTTCG 113
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 361 TCTTTTGTAAATTGGTGCACATATAGAGGTGAACAAGCTTATTAAAGGAAGATCCTTG 302

QY 114 CAGAAACATGCGCTAGTATCATTTGATGACCAACAT 147
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 301 TACATCAGAGTTTCATACANTAGATGAACGAT 268

RESULT 13
US-09-925-065A-691107/c
; Sequence 691107, Application US/09925065A
; Publication No. US20050228172A9
; GENERAL INFORMATION:
; APPLICANT: Wang, David G.
; TITLE OF INVENTION: Identification and Mapping of Single
; FILE OF INVENTION: Nucleotide Polymorphisms in the Human Genome
; FILE REFERENCE: 108827.135
; CURRENT APPLICATION NUMBER: US/09/925,065A
; CURRENT FILING DATE: 2001-08-08
; PRIOR APPLICATION NUMBER: US 60/243,096
; PRIOR FILING DATE: 2000-10-24
; PRIOR FILING DATE: 2000-11-20
; PRIOR FILING DATE: 2000-11-30
; PRIOR FILING DATE: 2001-01-16
; PRIOR APPLICATION NUMBER: US 60/289,846
; PRIOR FILING DATE: 2001-05-09
; NUMBER OF SEQ ID NOS: 957086
; SOFTWARE: FastSeq for Windows Version 4.0
; SEQ ID NO 691107
; LENGTH: 1437
; TYPE: DNA
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: misc feature
; LOCATION: 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18,
; LOCATION: 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32,
; LOCATION: 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46,
; LOCATION: 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60,
; LOCATION: 61, 62
; OTHER INFORMATION: n = A,T,C or G
```

```
; PRIOR APPLICATION NUMBER: US 60/261,766
; PRIOR FILING DATE: 2001-01-16
; PRIOR APPLICATION NUMBER: US 60/289,846
; PRIOR FILING DATE: 2001-05-09
; NUMBER OF SEQ ID NOS: 957086
; SOFTWARE: FastSeq for Windows Version 4.0
; SEQ ID NO 691107
; LENGTH: 1437
; TYPE: DNA
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: misc feature
; LOCATION: 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18,
; LOCATION: 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32,
; LOCATION: 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46,
; LOCATION: 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60,
; LOCATION: 61, 62
; OTHER INFORMATION: n = A,T,C or G
;
; NAME/KEY: misc feature
; LOCATION: 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77,
; LOCATION: 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91,
; LOCATION: 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102
; OTHER INFORMATION: n = A,T,C or G
US-09-925-065A-691107

Query Match 16.5%; Score 31.6; DB 4; Length 1437;
Best Local Similarity 55.5%; Pred. No. 11;
Matches 61; Conservative 0; Mismatches 49; Indels 0; Gaps 0;

QY 53 GTCTATTTTCATACCTTCGGGGAATAGATGTGAAACCCCTTATAAACCGCGGTTTTTC 112
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 479 GTCTTTGTGAAAGCCAAATGTGATATACATGTGAAAGCATTTTGTGNACTAAAGTGT 420

QY 113 GCAGAAACATGCGCTAGTATCATTTGATGACCAACATGCTCAATGCTGATGATGAAGAGGT 162
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 419 GTACAAACCAACCTAAATGTCCATCAATGCTGATGATGAAGAGGT 370

RESULT 14
US-09-925-065A-691108/c
; Sequence 691108, Application US/09925065A
; Publication No. US20050228172A9
; GENERAL INFORMATION:
; APPLICANT: Wang, David G.
; TITLE OF INVENTION: Identification and Mapping of Single
; FILE OF INVENTION: Nucleotide Polymorphisms in the Human Genome
; FILE REFERENCE: 108827.135
; CURRENT APPLICATION NUMBER: US/09/925,065A
; CURRENT FILING DATE: 2001-08-08
; PRIOR APPLICATION NUMBER: US 60/243,096
; PRIOR FILING DATE: 2000-10-24
; PRIOR FILING DATE: 2000-11-20
; PRIOR FILING DATE: 2000-11-30
; PRIOR FILING DATE: 2001-01-16
; PRIOR APPLICATION NUMBER: US 60/289,846
; PRIOR FILING DATE: 2001-05-09
; NUMBER OF SEQ ID NOS: 957086
; SOFTWARE: FastSeq for Windows Version 4.0
; SEQ ID NO 691108
; LENGTH: 1437
; TYPE: DNA
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: misc feature
; LOCATION: 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18,
; LOCATION: 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32,
; LOCATION: 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46,
; LOCATION: 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60,
; LOCATION: 61, 62
; OTHER INFORMATION: n = A,T,C or G
```

; OTHER INFORMATION: n = A,T,C or G  
; FEATURE:  
; NAME/KEY: misc.feature  
; LOCATION: 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77,  
; LOCATION: 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91,  
; LOCATION: 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102  
; OTHER INFORMATION: n = A,T,C or G  
US-09-925-065A-691108

Query Match 16.5%; Score 31.6; DB 4; Length 1437;  
Best Local Similarity 55.5%; Pred. No. 11;  
Matches 61; Conservative 0; Mismatches 49; Indels 0; Gaps 0;  
  
QY 53 GTCATTATTCATACCTTCGGGGAATAGATGTGAAAACCTTATATAAGCGGGTTTC 112  
|||||  
Db 479 GTCCTTGTGAAGCCAAATGTGATAATACATGTGAAGCATTTTGTGAACATAAAGTGT 420  
|||||  
  
QY 113 GCAGAAACATGCGCTAGTATCATTTGATGACAAATGGACTTAAGCAAAAGT 162  
|||||  
Db 419 GTACAAACACCTAAATGTCCATCATGGCTGAATGATAAAGAGGT 370  
|||||

RESULT 15  
US-10-674-124A-12097/c  
; Sequence 12097, Application US/10674124A  
; Publication No. US20040197797A1  
; GENERAL INFORMATION:  
; APPLICANT: INOKO, Hidetoshi  
; APPLICANT: TAMIYA, Gen  
; TITLE OF INVENTION: GENE MAPPING METHOD USING MICROSATELLITE  
; TITLE OF INVENTION: GENETIC POLYMORPHISM MARKERS  
; FILE REFERENCE: ORIN-003CIP  
; CURRENT APPLICATION NUMBER: US/10/674,124A  
; CURRENT FILING DATE: 2003-09-26  
; PRIOR APPLICATION NUMBER: 10/257,511  
; PRIOR FILING DATE: 2003-03-07  
; PRIOR APPLICATION NUMBER: PCT/JP00/07621  
; PRIOR FILING DATE: 2000-10-30  
; PRIOR APPLICATION NUMBER: JF2000-112699  
; PRIOR FILING DATE: 2000-04-13  
; PRIOR APPLICATION NUMBER: JF2002-327516  
; PRIOR FILING DATE: 2002-09-28  
; PRIOR APPLICATION NUMBER: JP2002-383869  
; PRIOR FILING DATE: 2002-12-09  
; NUMBER OF SEQ ID NOS: 27110  
; SEQ ID NO 12097  
; LENGTH: 360  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
; FEATURE:  
; OTHER INFORMATION: AC007735.2\_88547  
; FEATURE:  
; OTHER INFORMATION: Located on chromosome 7  
; FEATURE:  
; OTHER INFORMATION: Distance between a terminus base of telomere on  
; OTHER INFORMATION: chromosomal short arm and 5'-terminus of this base  
; OTHER INFORMATION: sequence : 54730897  
; FEATURE:  
; OTHER INFORMATION: Distance between 3'-terminus of neighbour sequence of  
; OTHER INFORMATION: sequence listing upward to telomere on chromosomal short arm and  
; OTHER INFORMATION: 5'-terminus of this base sequence : 7491  
US-10-674-124A-12097

Query Match 16.2%; Score 31.2; DB 8; Length 360;  
Best Local Similarity 54.3%; Pred. No. 7.9;  
Matches 63; Conservative 0; Mismatches 53; Indels 0; Gaps 0;  
  
QY 47 TGATTGCTCTATTTTCATACCTTCGGGGAATAGATGTGAAAACCTTATAAAACGCG 106  
|||||  
Db 313 TGCTTCATTTTTCATTAACCTTAAGAGAAATAGTATTCATTCCTTTGGTGAATTCGA 254  
|||||  
  
QY 107 GTTTTCGAGAAACATCGCTAGTATCATTTGATGACAACTGGACTAAGCAAAAGT 162  
|||||

Db 253 TGTCTACCAAAATGTGTCTGTCTAGTACTGATTACAGATTATAAAAAATAAAGT 198  
  
Search completed: March 7, 2006, 13:34:30  
Job time : 534.953 secs

GenCore version 5.1.7  
Copyright (c) 1993 - 2006 Bioceleration Ltd.

OM nucleic - nucleic search, using sw model

Run on: March 7, 2006, 12:18:50 ; Search time 1775.25 Seconds  
(without alignments)  
237.125 Million cell updates/sec

Title: US-10-782-899-1\_COPY\_1\_192

Perfect score: 192

Sequence: 1 gctggcgccggcgccatga.....tgaccacaagaagatgcttt 192

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 7218535 seqs, 1096242582 residues

Total number of hits satisfying chosen parameters: 14437070

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

- Database : Published Applications\_NA\_New.\*
- 1: /cgn2\_6/ptodata/1/pubpna/US08\_NEW\_PUB.seq.\*
  - 2: /cgn2\_6/ptodata/1/pubpna/US06\_NEW\_PUB.seq.\*
  - 3: /cgn2\_6/ptodata/1/pubpna/US07\_NEW\_PUB.seq.\*
  - 4: /cgn2\_6/ptodata/1/pubpna/PCT\_NEW\_PUB.seq.\*
  - 5: /cgn2\_6/ptodata/1/pubpna/US09\_NEW\_PUB.seq.\*
  - 6: /cgn2\_6/ptodata/1/pubpna/US09\_NEW\_PUB.seq1.\*
  - 7: /cgn2\_6/ptodata/1/pubpna/US10\_NEW\_PUB.seq.\*
  - 8: /cgn2\_6/ptodata/1/pubpna/US10\_NEW\_PUB.seq1.\*
  - 9: /cgn2\_6/ptodata/1/pubpna/US11\_NEW\_PUB.seq.\*
  - 10: /cgn2\_6/ptodata/1/pubpna/US11\_NEW\_PUB.seq2.\*
  - 11: /cgn2\_6/ptodata/1/pubpna/US11\_NEW\_PUB.seq3.\*
  - 12: /cgn2\_6/ptodata/1/pubpna/US11\_NEW\_PUB.seq4.\*
  - 13: /cgn2\_6/ptodata/1/pubpna/US60\_NEW\_PUB.seq.\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
C 1	31.6	16.5	1437	6	US-09-925-065A-691107 Sequence 691107,
C 2	31.6	16.5	1437	6	US-09-925-065A-691108 Sequence 691108,
C 3	31.2	16.2	1485	6	US-09-925-065A-49312 Sequence 49312, A
C 4	31.2	16.2	1485	6	US-09-925-065A-49313 Sequence 49313, A
C 5	31.2	16.2	96109	12	US-11-124-368A-2895 Sequence 2895, Ap
C 6	31	16.1	1771	6	US-09-925-065A-65588 Sequence 65588, A
C 7	30.8	16.0	639	6	US-09-925-065A-170277 Sequence 170277,
C 8	30.8	16.0	639	6	US-09-925-065A-170278 Sequence 170278,
C 9	30.6	15.9	572	6	US-09-925-065A-308556 Sequence 308556,
C 10	30.6	15.9	572	6	US-09-925-065A-308557 Sequence 308557,
C 11	30.6	15.9	601	6	US-09-925-065A-396377 Sequence 396377,
C 12	30.6	15.9	606	6	US-09-925-065A-598874 Sequence 598874,
C 13	30.6	15.9	670	6	US-09-925-065A-905740 Sequence 905740,
C 14	30.6	15.9	670	6	US-09-925-065A-912520 Sequence 912520,
C 15	30.4	15.8	528	6	US-09-925-065A-149916 Sequence 149916,
C 16	30.4	15.8	774	8	US-10-750-185-44581 Sequence 44581, A
C 17	30.4	15.8	774	8	US-10-750-623-44581 Sequence 44581, A
C 18	30.4	15.8	1569	8	US-10-793-626-603 Sequence 603, App
C 19	30.4	15.8	3204	8	US-10-793-626-3485 Sequence 3485, Ap
C 20	30.2	15.7	583	6	US-09-925-065A-460994 Sequence 460994,

ALIGNMENTS

RESULT 1

US-09-925-065A-691107/c  
; Sequence 691107, Application US/09925065A  
; Publication No. US20040181048A1  
; GENERAL INFORMATION:  
; APPLICANT: Wang, David G.  
; TITLE OF INVENTION: Identification and Mapping of Single  
; FILE REFERENCE: Nucleotide Polymorphisms in the Human Genome  
; CURRENT APPLICATION NUMBER: US/09/925,065A  
; PRIOR FILING DATE: 2001-08-08  
; PRIOR APPLICATION NUMBER: US 60/243,096  
; PRIOR FILING DATE: 2000-10-24  
; PRIOR APPLICATION NUMBER: US 60/252,147  
; PRIOR FILING DATE: 2000-11-20  
; PRIOR APPLICATION NUMBER: US 60/250,092  
; PRIOR FILING DATE: 2000-11-30  
; PRIOR APPLICATION NUMBER: US 60/261,766  
; PRIOR FILING DATE: 2001-01-16  
; PRIOR APPLICATION NUMBER: US 60/289,846  
; PRIOR FILING DATE: 2001-05-09  
; NUMBER OF SEQ ID NOS: 957086  
; SOFTWARE: FastSeq for Windows Version 4.0  
; SEQ ID NO 691107  
; LENGTH: 1437  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
; FEATURE:  
; NAME/KEY: misc\_feature  
; LOCATION: 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18,  
; LOCATION: 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32,  
; LOCATION: 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46,  
; LOCATION: 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60,  
; LOCATION: 61, 62  
; OTHER INFORMATION: n = A,T,C or G  
; FEATURE:  
; NAME/KEY: misc\_feature  
; LOCATION: 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77,  
; LOCATION: 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91,  
; LOCATION: 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102  
; OTHER INFORMATION: n = A,T,C or G  
US-09-925-065A-691107

Sequence 684507,  
Sequence 684508,  
Sequence 13302, A  
Sequence 51223, A  
Sequence 463323,  
Sequence 355910,  
Sequence 157, App  
Sequence 160, App  
Sequence 140508,  
Sequence 398346,  
Sequence 398347,  
Sequence 509858,  
Sequence 5015, Ap  
Sequence 795197,  
Sequence 797051,  
Sequence 616829,  
Sequence 616830,  
Sequence 616831,  
Sequence 616832,  
Sequence 795196,  
Sequence 613398,  
Sequence 45, Appl  
Sequence 150, App  
Sequence 83, Appl  
Sequence 148, App

Query Match 16.5%; Score 31.6; DB 6; Length 1437;  
Best Local Similarity 55.5%; Pred. No. 9.4;  
Matches 61; Conservative 0; Mismatches 49; Indels 0; Gaps 0;  
  
QY 53 GTCATTTTCAATACCTTCGGGAAATAGATGTGAAACCCCTTATAAAACGCGGTTTC 112  
|||||  
Db 479 GTCCTTGTGAAGCCAAATGTGATATACATGTGAAAGCATTTTGTGAACCTAAAGTGT 420  
|||||  
  
QY 113 GCAGAACATGCCTAGTATCATTTGATGACCAATGGAATGAAAGT 162  
|||||  
Db 419 GTACAAACACCTAAATGTCCATCAATGGCTGAATGATTAAGAGAGGT 370  
|||||

RESULT 2  
US-09-925-065A-691108/c  
; Sequence 691108, Application US/09925065A  
; Publication No. US20040181048A1  
; GENERAL INFORMATION:  
; APPLICANT: Wang, David G.  
; TITLE OF INVENTION: Nucleotide Polymorphisms in the Human Genome  
; FILE REFERENCE: 108827.135  
; CURRENT APPLICATION NUMBER: US 60/243,096  
; PRIOR FILING DATE: 2001-08-08  
; PRIOR APPLICATION NUMBER: US 60/252,147  
; PRIOR FILING DATE: 2000-10-24  
; PRIOR FILING DATE: 2000-11-20  
; PRIOR APPLICATION NUMBER: US 60/250,092  
; PRIOR FILING DATE: 2000-11-30  
; PRIOR APPLICATION NUMBER: US 60/261,766  
; PRIOR FILING DATE: 2001-01-16  
; PRIOR APPLICATION NUMBER: US 60/289,846  
; PRIOR FILING DATE: 2001-05-09  
; NUMBER OF SEQ ID NOS: 957086  
; SOFTWARE: FastSeq for Windows Version 4.0  
; SEQ ID NO 691108  
; LENGTH: 1437  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
; FEATURE:  
; NAME/KEY: misc feature  
; LOCATION: 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18,  
; LOCATION: 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32,  
; LOCATION: 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46,  
; LOCATION: 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60,  
; LOCATION: 61, 62  
; OTHER INFORMATION: n = A,T,C or G  
; FEATURE:  
; NAME/KEY: misc feature  
; LOCATION: 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77,  
; LOCATION: 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91,  
; LOCATION: 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102  
; OTHER INFORMATION: n = A,T,C or G  
US-09-925-065A-691108

Query Match 16.5%; Score 31.6; DB 6; Length 1437;  
Best Local Similarity 55.5%; Pred. No. 9.4;  
Matches 61; Conservative 0; Mismatches 49; Indels 0; Gaps 0;  
  
QY 53 GTCATTTTCAATACCTTCGGGAAATAGATGTGAAACCCCTTATAAAACGCGGTTTC 112  
|||||  
Db 479 GTCCTTGTGAAGCCAAATGTGATATACATGTGAAAGCATTTTGTGAACCTAAAGTGT 420  
|||||  
  
QY 113 GCAGAACATGCCTAGTATCATTTGATGACCAATGGAATGAAAGT 162  
|||||  
Db 419 GTACAAACACCTAAATGTCCATCAATGGCTGAATGATTAAGAGAGGT 370  
|||||

RESULT 3  
US-09-925-065A-49312  
; Sequence 49312, Application US/09925065A  
; Publication No. US20040181048A1

GENERAL INFORMATION:  
; APPLICANT: Wang, David G.  
; TITLE OF INVENTION: Identification and Mapping of Single  
; FILE REFERENCE: 108827.135  
; CURRENT APPLICATION NUMBER: US/09/925,065A  
; CURRENT FILING DATE: 2001-08-08  
; PRIOR APPLICATION NUMBER: US 60/243,096  
; PRIOR FILING DATE: 2000-10-24  
; PRIOR APPLICATION NUMBER: US 60/252,147  
; PRIOR FILING DATE: 2000-11-20  
; PRIOR APPLICATION NUMBER: US 60/250,092  
; PRIOR FILING DATE: 2000-11-30  
; PRIOR APPLICATION NUMBER: US 60/261,766  
; PRIOR FILING DATE: 2001-01-16  
; PRIOR APPLICATION NUMBER: US 60/289,846  
; PRIOR FILING DATE: 2001-05-09  
; NUMBER OF SEQ ID NOS: 957086  
; SOFTWARE: FastSeq for Windows Version 4.0  
; SEQ ID NO 49312  
; LENGTH: 1485  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
US-09-925-065A-49312

Query Match 16.2%; Score 31.2; DB 6; Length 1485;  
Best Local Similarity 57.0%; Pred. No. 12;  
Matches 57; Conservative 0; Mismatches 43; Indels 0; Gaps 0;  
  
QY 20 AAGTGGCTTGACAAGCATATCTTGCTGATTCGTCTATTTTCAATACCTTCGGGAAAT 79  
|||||  
Db 1062 AAGTCAATGCTAACCATTTCTTCTATTTTCTTATATTTCTTATAGCTTAGGTGACAT 1121  
|||||  
  
QY 80 AGATGTGAAACCCCTTATAAAACGCGGTTTCGCGAGAA 119  
|||||  
Db 1122 AATAGGAAGATGTTTGTACTCTAGCATAATAAATAAAA 1161  
|||||

RESULT 4  
US-09-925-065A-49313  
; Sequence 49313, Application US/09925065A  
; Publication No. US20040181048A1  
; GENERAL INFORMATION:  
; APPLICANT: Wang, David G.  
; TITLE OF INVENTION: Identification and Mapping of Single  
; FILE REFERENCE: 108827.135  
; CURRENT APPLICATION NUMBER: US/09/925,065A  
; CURRENT FILING DATE: 2001-08-08  
; PRIOR APPLICATION NUMBER: US 60/243,096  
; PRIOR FILING DATE: 2000-10-24  
; PRIOR APPLICATION NUMBER: US 60/252,147  
; PRIOR FILING DATE: 2000-11-20  
; PRIOR APPLICATION NUMBER: US 60/250,092  
; PRIOR FILING DATE: 2000-11-30  
; PRIOR APPLICATION NUMBER: US 60/261,766  
; PRIOR FILING DATE: 2001-01-16  
; PRIOR APPLICATION NUMBER: US 60/289,846  
; PRIOR FILING DATE: 2001-05-09  
; NUMBER OF SEQ ID NOS: 957086  
; SOFTWARE: FastSeq for Windows Version 4.0  
; SEQ ID NO 49313  
; LENGTH: 1485  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
US-09-925-065A-49313

Query Match 16.2%; Score 31.2; DB 6; Length 1485;  
Best Local Similarity 57.0%; Pred. No. 12;  
Matches 57; Conservative 0; Mismatches 43; Indels 0; Gaps 0;  
  
QY 20 AAGTGGCTTGACAAGCATATCTTGCTGATTCGTCTATTTTCAATACCTTCGGGAAAT 79  
|||||

Db 1062 AAGTCATCTGCTAACCAATTTCTTCTATTTCTTATATTTCTTATATTTCTTATAGCTTAGGTGACAT 11121  
QY 80 AGATGTGAACACCTTATAAAGCGGGTTTTTCGCAGAAA 119  
Db 1122 AAATAGGAGATGTTTGTACTCTAGCATATAAATAAAA 1161

## RESULT 5

US-11-124-368A-2895/c  
; Sequence 2895, Application US/11124368A  
; Publication No. US20050287559A1  
; GENERAL INFORMATION:  
; APPLICANT: Michele Cargill  
; APPLICANT: James J. Devlin  
; TITLE OF INVENTION: Genetic Polymorphisms Associated with  
; TITLE OF INVENTION: Vascular Diseases, Methods of Detection and Uses Thereof  
; FILE REFERENCE: CL001524  
; CURRENT APPLICATION NUMBER: US/11/124,368A  
; CURRENT FILING DATE: 2005-05-09  
; PRIOR APPLICATION NUMBER: US 60/568,845  
; PRIOR FILING DATE: 2004-05-07  
; PRIOR APPLICATION NUMBER: US 60/625,936  
; PRIOR FILING DATE: 2004-11-09  
; NUMBER OF SEQ ID NOS: 21112  
; SOFTWARE: FastSeq for Windows Version 4.0  
; SEQ ID NO 2895  
; LENGTH: 96109  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
US-11-124-368A-2895

Query Match 16.2%; Score 31.2; DB 12; Length 96109;  
Best Local Similarity 53.2%; Pred. No. 43;  
Matches 66; Conservative 0; Mismatches 58; Indels 0; Gaps 0;  
QY 63 AATACTTCGGGGAATAGATGTGAACACCTTATAAAGCGGGTTTTTCGCAGAACAT 122  
Db 63419 AAGGACCTCAAGGAGTATAGAAAAACCTTCCCAAGCTGGATGATGAAGGAGAT 63360  
QY 123 GCCTAGTATCATTTGATGACACATGACCTAAGCAAAAGTCTTCCCTGACCCCAAGA 182  
Db 63359 GAGACAGTGACAGAGAGACGAGGGGCTCTCATATTTGTGTGTAGTACCAACT 63300  
QY 183 AGGA 186  
Db 63299 AAGA 63296

## RESULT 6

US-09-925-065A-65588  
; Sequence 65588, Application US/09925065A  
; Publication No. US20040181048A1  
; GENERAL INFORMATION:  
; APPLICANT: Wang, David G.  
; TITLE OF INVENTION: Identification and Mapping of Single  
; TITLE OF INVENTION: Nucleotide Polymorphisms in the Human Genome  
; FILE REFERENCE: 108827.135  
; CURRENT APPLICATION NUMBER: US/09/925,065A  
; CURRENT FILING DATE: 2001-08-08  
; PRIOR APPLICATION NUMBER: US 60/243,096  
; PRIOR FILING DATE: 2000-10-24  
; PRIOR APPLICATION NUMBER: US 60/252,147  
; PRIOR FILING DATE: 2000-11-20  
; PRIOR APPLICATION NUMBER: US 60/250,092  
; PRIOR FILING DATE: 2000-11-30  
; PRIOR APPLICATION NUMBER: US 60/261,766  
; PRIOR FILING DATE: 2001-01-16  
; PRIOR APPLICATION NUMBER: US 60/289,846  
; PRIOR FILING DATE: 2001-05-09  
; NUMBER OF SEQ ID NOS: 957086  
; SOFTWARE: FastSeq for Windows Version 4.0  
; SEQ ID NO 65588

; LENGTH: 1771  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
US-09-925-065A-65588

Query Match 16.1%; Score 31; DB 6; Length 1771;  
Best Local Similarity 53.8%; Pred. No. 15;  
Matches 64; Conservative 0; Mismatches 55; Indels 0; Gaps 0;  
QY 55 CTATTTTCAATACCTTCGGGGAATAGATGTGAACACCTTATAAAGCGGGTTTTTCGC 114  
Db 1186 CTAACCTAAATGCTTCAGGCAATAAATTGCAACTCATATGAATAGACAAATTCCT 1245  
QY 115 AGAACAATCGCTAGTATCATTTGATGACCAATGGAAGCAAAAGTCTTTCGCCCT 173  
Db 1246 AGAAAAACGCACATGAATAAATAATATATATATATATATATATATATATATATAT 1304

## RESULT 7

US-09-925-065A-170277/c  
; Sequence 170277, Application US/09925065A  
; Publication No. US20040181048A1  
; GENERAL INFORMATION:  
; APPLICANT: Wang, David G.  
; TITLE OF INVENTION: Identification and Mapping of Single  
; TITLE OF INVENTION: Nucleotide Polymorphisms in the Human Genome  
; FILE REFERENCE: 108827.135  
; CURRENT APPLICATION NUMBER: US/09/925,065A  
; CURRENT FILING DATE: 2001-08-08  
; PRIOR APPLICATION NUMBER: US 60/243,096  
; PRIOR FILING DATE: 2000-10-24  
; PRIOR APPLICATION NUMBER: US 60/252,147  
; PRIOR FILING DATE: 2000-11-20  
; PRIOR APPLICATION NUMBER: US 60/250,092  
; PRIOR FILING DATE: 2000-11-30  
; PRIOR APPLICATION NUMBER: US 60/261,766  
; PRIOR FILING DATE: 2001-01-16  
; PRIOR APPLICATION NUMBER: US 60/289,846  
; PRIOR FILING DATE: 2001-05-09  
; NUMBER OF SEQ ID NOS: 957086  
; SOFTWARE: FastSeq for Windows Version 4.0  
; SEQ ID NO 170277  
; LENGTH: 639  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
US-09-925-065A-170277

Query Match 16.0%; Score 30.8; DB 6; Length 639;  
Best Local Similarity 55.7%; Pred. No. 13;  
Matches 59; Conservative 0; Mismatches 47; Indels 0; Gaps 0;  
QY 43 TGCTGATTCGTCTATTTTCAATACCTTCGGGGAATAGATGTGAACACCTTATAAAG 102  
Db 616 TTCTGATTTGTGTAATTTTCTTCTCTGAAAAATTTAGTTTACCATATCTTGTAT 557  
QY 103 GCGGGTTTTTCGCAGAACATGCGCTAGTATCATTTGATGACCAATG 148  
Db 556 GCAGATTTCTGGAGCAATTCCTGTTTTTGTATATCTCAAAAG 511

## RESULT 8

US-09-925-065A-170278/c  
; Sequence 170278, Application US/09925065A  
; Publication No. US20040181048A1  
; GENERAL INFORMATION:  
; APPLICANT: Wang, David G.  
; TITLE OF INVENTION: Identification and Mapping of Single  
; TITLE OF INVENTION: Nucleotide Polymorphisms in the Human Genome  
; FILE REFERENCE: 108827.135  
; CURRENT APPLICATION NUMBER: US/09/925,065A  
; CURRENT FILING DATE: 2001-08-08  
; PRIOR APPLICATION NUMBER: US 60/243,096  
; PRIOR FILING DATE: 2000-10-24

; PRIOR APPLICATION NUMBER: US 60/252,147  
; PRIOR FILING DATE: 2000-11-20  
; PRIOR APPLICATION NUMBER: US 60/250,092  
; PRIOR FILING DATE: 2000-11-30  
; PRIOR APPLICATION NUMBER: US 60/261,766  
; PRIOR FILING DATE: 2001-01-16  
; PRIOR APPLICATION NUMBER: US 60/289,846  
; PRIOR FILING DATE: 2001-05-09  
; NUMBER OF SEQ ID NOS: 957086  
; SOFTWARE: FastSEQ for Windows Version 4.0  
; SEQ ID NO 170278  
; LENGTH: 639  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
US-09-925-065A-170278

Query Match 16.0%; Score 30.8; DB 6; Length 639;  
Best Local Similarity 55.7%; Pred. No. 13;  
Matches 59; Conservative 0; Mismatches 47; Indels 0; Gaps 0;  
QY 43 TGCTGATTCGCTATTTTCAATACCTTCGGGAAATAGATGTGAAAACCCITTATAAAC 102  
DB 616 TTTCTGATTTGTGTAATTTTCTTCTGAAAAATTTAGTTTATACCATCTTGTAAAT 557  
QY 103 GCGGGTTTTCGAGAACATCGCTAGTATCATTTGATGACCAACATG 148  
DB 556 GCAGATTTCTGGAGACAAATCTCTGTTTGTGTTTATCTCAAAAAG 511

RESULT 9  
US-09-925-065A-308556/c  
; Sequence 308556, Application US/09925065A  
; Publication No. US20040181048A1  
; GENERAL INFORMATION:  
; APPLICANT: Wang, David G.  
; TITLE OF INVENTION: Identification and Mapping of Single  
; FILE REFERENCE: 108827.135  
; CURRENT APPLICATION NUMBER: US/09/925,065A  
; CURRENT FILING DATE: 2001-08-08  
; PRIOR APPLICATION NUMBER: US 60/243,096  
; PRIOR FILING DATE: 2000-10-24  
; PRIOR APPLICATION NUMBER: US 60/252,147  
; PRIOR FILING DATE: 2000-11-20  
; PRIOR APPLICATION NUMBER: US 60/250,092  
; PRIOR FILING DATE: 2000-11-30  
; PRIOR APPLICATION NUMBER: US 60/261,766  
; PRIOR FILING DATE: 2001-01-16  
; PRIOR APPLICATION NUMBER: US 60/289,846  
; PRIOR FILING DATE: 2001-05-09  
; NUMBER OF SEQ ID NOS: 957086  
; SOFTWARE: FastSEQ for Windows Version 4.0  
; SEQ ID NO 308556  
; LENGTH: 572  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
US-09-925-065A-308556

Query Match 15.9%; Score 30.6; DB 6; Length 572;  
Best Local Similarity 50.3%; Pred. No. 14;  
Matches 75; Conservative 0; Mismatches 74; Indels 0; Gaps 0;  
QY 35 CATAACTTCTGATTCGCTATTTTCAATACCTTCGGGAAATAGATGTGAAAACCCCT 94  
DB 397 CATGGTTTATTTAAATTTGAGACTTTTCAAAACCTTGTGTGTTTATATACCAACATA 338  
QY 95 TATAAAACGGGGTTTTCGAGAAACATGCGCTAGTATCATTTGATGACAAACATGGACTAA 154  
DB 337 AATTGAAAACAGCTTTTACCAGACATTTAAGATAGCCATGATTAAATAACATGAAAT 278  
QY 155 GCAAAAGTGTCTGTCCCTGACCCAGAA 183  
DB 277 TTAATAATGTTTGCCTATTTACCCATGTA 249

RESULT 10  
US-09-925-065A-308557/c  
; Sequence 308557, Application US/09925065A  
; Publication No. US20040181048A1  
; GENERAL INFORMATION:  
; APPLICANT: Wang, David G.  
; TITLE OF INVENTION: Identification and Mapping of Single  
; FILE REFERENCE: 108827.135  
; CURRENT APPLICATION NUMBER: US/09/925,065A  
; CURRENT FILING DATE: 2001-08-08  
; PRIOR APPLICATION NUMBER: US 60/243,096  
; PRIOR FILING DATE: 2000-10-24  
; PRIOR APPLICATION NUMBER: US 60/252,147  
; PRIOR FILING DATE: 2000-11-20  
; PRIOR APPLICATION NUMBER: US 60/250,092  
; PRIOR FILING DATE: 2000-11-30  
; PRIOR APPLICATION NUMBER: US 60/261,766  
; PRIOR FILING DATE: 2001-01-16  
; PRIOR APPLICATION NUMBER: US 60/289,846  
; PRIOR FILING DATE: 2001-05-09  
; NUMBER OF SEQ ID NOS: 957086  
; SOFTWARE: FastSEQ for Windows Version 4.0  
; SEQ ID NO 308557  
; LENGTH: 572  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
US-09-925-065A-308557

Query Match 15.9%; Score 30.6; DB 6; Length 572;  
Best Local Similarity 50.3%; Pred. No. 14;  
Matches 75; Conservative 0; Mismatches 74; Indels 0; Gaps 0;  
QY 35 CATAACTTCTGATTCGCTATTTTCAATACCTTCGGGAAATAGATGTGAAAACCCCT 94  
DB 397 CATGGTTTATTTAAATTTGAGACTTTTCAAAACCTTGTGTGTTTATATACCAACATA 338  
QY 95 TATAAAACGGGGTTTTCGAGAAACATGCGCTAGTATCATTTGATGACAAACATGGACTAA 154  
DB 337 AATTGAAAACAGCTTTTACCAGACATTTAAGATAGCCATGATTAAATAACATGAAAT 278  
QY 155 GCAAAAGTGTCTGTCCCTGACCCAGAA 183  
DB 277 TTAATAATGTTTGCCTATTTACCCATGTA 249

RESULT 11  
US-09-925-065A-396377/c  
; Sequence 396377, Application US/09925065A  
; Publication No. US20040181048A1  
; GENERAL INFORMATION:  
; APPLICANT: Wang, David G.  
; TITLE OF INVENTION: Identification and Mapping of Single  
; FILE REFERENCE: 108827.135  
; CURRENT APPLICATION NUMBER: US/09/925,065A  
; CURRENT FILING DATE: 2001-08-08  
; PRIOR APPLICATION NUMBER: US 60/243,096  
; PRIOR FILING DATE: 2000-10-24  
; PRIOR APPLICATION NUMBER: US 60/252,147  
; PRIOR FILING DATE: 2000-11-20  
; PRIOR APPLICATION NUMBER: US 60/250,092  
; PRIOR FILING DATE: 2000-11-30  
; PRIOR APPLICATION NUMBER: US 60/261,766  
; PRIOR FILING DATE: 2001-01-16  
; PRIOR APPLICATION NUMBER: US 60/289,846  
; PRIOR FILING DATE: 2001-05-09  
; NUMBER OF SEQ ID NOS: 957086  
; SOFTWARE: FastSEQ for Windows Version 4.0  
; SEQ ID NO 396377  
; LENGTH: 601



```
; TYPE: DNA
; ORGANISM: Homo sapiens
US-09-925-065A-396377

Query Match      15.9%; Score 30.6; DB 6; Length 601;
Best Local Similarity 58.1%; Pred. No. 15;
Matches 54; Conservative 0; Mismatches 39; Indels 0; Gaps 0;

QY 56 TATTTCATACCTTCGGGGAATAGATGTGAAACCCCTTATATAACGGCGGTTTTCGCA 115
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 544 TACTTTCATTAACCTTAGGAAGCTGAAGGGGAAAAATCAATAAGCAGCTCTCTTGACT 485

QY 116 GAAACATGCGCTAGTATCATTTGATGACAACATG 148
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 484 CAAAGATGAGCTGAAATAATACATGAATGAATG 452

RESULT 12
US-09-925-065A-599874
; Sequence 599874, Application US/09925065A
; Publication No. US20040181048A1
; GENERAL INFORMATION:
; APPLICANT: Wang, David G.
; TITLE OF INVENTION: Identification and Mapping of Single
; FILE OF INVENTION: Nucleotide Polymorphisms in the Human Genome
; FILE REFERENCE: 108827.135
; CURRENT APPLICATION NUMBER: US/09/925,065A
; CURRENT FILING DATE: 2001-08-08
; PRIOR APPLICATION NUMBER: US 60/243,096
; PRIOR FILING DATE: 2000-10-24
; PRIOR APPLICATION NUMBER: US 60/252,147
; PRIOR FILING DATE: 2000-11-20
; PRIOR APPLICATION NUMBER: US 60/250,092
; PRIOR FILING DATE: 2000-11-30
; PRIOR APPLICATION NUMBER: US 60/261,766
; PRIOR FILING DATE: 2001-01-16
; PRIOR APPLICATION NUMBER: US 60/289,846
; PRIOR FILING DATE: 2001-05-09
; NUMBER OF SEQ ID NOS: 957086
; SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO 599874
; LENGTH: 606
; TYPE: DNA
; ORGANISM: Homo sapiens
US-09-925-065A-599874

Query Match      15.9%; Score 30.6; DB 6; Length 606;
Best Local Similarity 58.1%; Pred. No. 15;
Matches 54; Conservative 0; Mismatches 39; Indels 0; Gaps 0;

QY 56 TATTTCATACCTTCGGGGAATAGATGTGAAACCCCTTATATAACGGCGGTTTTCGCA 115
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 63 TACTTTCATTAACCTTAGGAAGCTGAAGGGGAAAAATCAATAAGCAGCTCTCTTGACT 122

QY 116 GAAACATGCGCTAGTATCATTTGATGACAACATG 148
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 123 CAAAGATGAGCTGAAATAATACATGAATGAATG 155

RESULT 13
US-09-925-065A-905740
; Sequence 905740, Application US/09925065A
; Publication No. US20040181048A1
; GENERAL INFORMATION:
; APPLICANT: Wang, David G.
; TITLE OF INVENTION: Identification and Mapping of Single
; FILE OF INVENTION: Nucleotide Polymorphisms in the Human Genome
; FILE REFERENCE: 108827.135
; CURRENT APPLICATION NUMBER: US/09/925,065A
; CURRENT FILING DATE: 2001-08-08
; PRIOR APPLICATION NUMBER: US 60/243,096
; PRIOR FILING DATE: 2000-10-24
; PRIOR APPLICATION NUMBER: US 60/252,147
```

```
; PRIOR FILING DATE: 2000-11-20
; PRIOR APPLICATION NUMBER: US 60/250,092
; PRIOR FILING DATE: 2000-11-30
; PRIOR APPLICATION NUMBER: US 60/261,766
; PRIOR FILING DATE: 2001-01-16
; PRIOR APPLICATION NUMBER: US 60/289,846
; PRIOR FILING DATE: 2001-05-09
; NUMBER OF SEQ ID NOS: 957086
; SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO 905740
; LENGTH: 670
; TYPE: DNA
; ORGANISM: Homo sapiens
US-09-925-065A-905740

Query Match      15.9%; Score 30.6; DB 6; Length 670;
Best Local Similarity 60.0%; Pred. No. 15;
Matches 51; Conservative 0; Mismatches 34; Indels 0; Gaps 0;

QY 21 AGTGGCTTGACAAGCATATCTTGCTGATTCGTCTATTTTCAATACCTTCGGGGAAATA 80
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 541 ACTAACTAGACAGAGATATCTTGTAATTTACTTTTGTGTTTACATACACTACATATAATA 600

QY 81 GATGGAACCCCTTATAAACGCG 105
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 601 TATGATATAACATGTATAGACGAG 625

RESULT 14
US-09-925-065A-912520/c
; Sequence 912520, Application US/09925065A
; Publication No. US20040181048A1
; GENERAL INFORMATION:
; APPLICANT: Wang, David G.
; TITLE OF INVENTION: Identification and Mapping of Single
; FILE OF INVENTION: Nucleotide Polymorphisms in the Human Genome
; FILE REFERENCE: 108827.135
; CURRENT APPLICATION NUMBER: US/09/925,065A
; CURRENT FILING DATE: 2001-08-08
; PRIOR APPLICATION NUMBER: US 60/243,096
; PRIOR FILING DATE: 2000-10-24
; PRIOR APPLICATION NUMBER: US 60/252,147
; PRIOR FILING DATE: 2000-11-20
; PRIOR APPLICATION NUMBER: US 60/250,092
; PRIOR FILING DATE: 2000-11-30
; PRIOR APPLICATION NUMBER: US 60/261,766
; PRIOR FILING DATE: 2001-01-16
; PRIOR APPLICATION NUMBER: US 60/289,846
; PRIOR FILING DATE: 2001-05-09
; NUMBER OF SEQ ID NOS: 957086
; SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO 912520
; LENGTH: 670
; TYPE: DNA
; ORGANISM: Homo sapiens
US-09-925-065A-912520

Query Match      15.9%; Score 30.6; DB 6; Length 670;
Best Local Similarity 60.0%; Pred. No. 15;
Matches 51; Conservative 0; Mismatches 34; Indels 0; Gaps 0;

QY 21 AGTGGCTTGACAAGCATATCTTGCTGATTCGTCTATTTTCAATACCTTCGGGGAAATA 80
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 130 ACTAACTAGACAGAGATATCTTGTAATTTACTTTTGTGTTTACATACACTACATATAATA 71

QY 81 GATGGAACCCCTTATAAACGCG 105
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 70 TATGATATAACATGTATAGACGAG 46

RESULT 15
US-09-925-065A-149916
; Sequence 149916, Application US/09925065A
```

```
; Publication No. US20040181048A1
; GENERAL INFORMATION:
; APPLICANT: Wang, David G.
; TITLE OF INVENTION: Identification and Mapping of Single
; TITLE OF INVENTION: Nucleotide Polymorphisms in the Human Genome
; FILE REFERENCE: 108827.135
; CURRENT APPLICATION NUMBER: US/09/925,065A
; CURRENT FILING DATE: 2001-08-08
; PRIOR APPLICATION NUMBER: US 60/243,096
; PRIOR FILING DATE: 2000-10-24
; PRIOR APPLICATION NUMBER: US 60/252,147
; PRIOR FILING DATE: 2000-11-20
; PRIOR APPLICATION NUMBER: US 60/250,092
; PRIOR FILING DATE: 2000-11-30
; PRIOR APPLICATION NUMBER: US 60/261,766
; PRIOR FILING DATE: 2001-01-16
; PRIOR APPLICATION NUMBER: US 60/289,846
; PRIOR FILING DATE: 2001-05-09
; NUMBER OF SEQ ID NOS: 957086
; SOFTWARE: FastSeq for Windows Version 4.0
; SEQ ID NO 149916
; LENGTH: 528
; TYPE: DNA
; ORGANISM: Homo sapiens
; US-09-925-065A-149916

Query Match      15.8%; Score 30.4; DB 6; Length 528;
Best Local Similarity 50.0%; Pred. No. 16;
Matches 73; Conservative 1; Mismatches 72; Indels 0; Gaps 0;

QY 20 AAGTGGCTTGACAAAGCATATCTGTCTGATTCTGCTATTTCATACCTTCGGGAAAT 79
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 365 AAGTTTCTTAAAGAAATATTTCTGTTGTTGTTTTCATTACCTTACCTAAGTCGATA 424

QY 80 AGATGTGAAACCCCTTATATAAACCGGGGTTTTTCGAGAAACATGCGCTAGTATCATTTGAT 139
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 425 GAATGTAATCTTTCTTACATGATTGTTTAGTCTAATCAAAATGCTGAAGCATAAGTGAT 484

QY 140 GACAAACATGGACTAAGCAAAAGTGCT 165
    |: | | | | | | | | | |
Db 485 AAYCCTCCATAATGATACAAACTGAT 510
```

Search completed: March 7, 2006, 13:20:42  
Job time : 1776.25 secs

GenCore version 5.1.7  
Copyright (c) 1993 - 2006 Bioceleration Ltd.

OM nucleic - nucleic search, using sw model

Run on: March 7, 2006, 11:25:13 ; Search time 1196.77 Seconds  
(without alignments)  
6127.182 Million cell updates/sec

Title: US-10-782-899-1\_COPY\_472\_600

Perfect score: 129  
Sequence: 1 tgacctctcgtcgtagcga.....tattccgactgattcagcg 129

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 588141 seqs, 28421725653 residues

Total number of hits satisfying chosen parameters: 11766282

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

GenEmbl:\*

- 1: gb\_da:\*
- 2: gb\_in:\*
- 3: gb\_env:\*
- 4: gb\_sm:\*
- 5: gb\_ov:\*
- 6: gb\_pat:\*
- 7: gb\_ph:\*
- 8: gb\_pr:\*
- 9: gb\_ro:\*
- 10: gb\_sts:\*
- 11: gb\_sv:\*
- 12: gb\_un:\*
- 13: gb\_vi:\*
- 14: gb\_htg:\*
- 15: gb\_pl:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	129	100.0	600	1	AB072446
2	129	100.0	600	6	BD144660
C 3	129	100.0	56640	1	AE017283_21
C 4	129	100.0	156638	6	AX492788
C 5	129	100.0	156646	6	AX533955
C 6	32.6	25.3	295650	1	BX294151
C 7	32	24.8	194439	2	CNS07EGA
C 8	31.8	24.7	3048	15	AK119844
C 9	31.4	24.3	90417	14	AC138542
C 10	31.4	24.3	92770	14	AC138544
C 11	31.4	24.3	128428	14	AC144889
C 12	31.4	24.3	207821	14	AC093397
C 13	31	24.0	5832	1	AF105113
C 14	31	24.0	18617	1	CR931675
C 15	31	24.0	18754	1	AF094575
C 16	30.8	23.9	222330	14	AC096255
C 17	30.8	23.9	246369	14	AC125707
C 18	30.8	23.9	302050	1	AL935257

19	30.6	23.7	821	10	BV633898
20	30.6	23.7	26309	6	CQ363747
21	30.6	23.7	110000	1	AE017283_21
22	30.6	23.7	223086	14	AC130043
23	30.6	23.7	229616	14	AC156070
C 24	30	23.3	960	1	AY343890
25	30	23.3	236921	14	AC094151
C 26	29.8	23.1	100326	9	AL669928
27	29.8	23.1	110000	15	AP008214_161
28	29.8	23.1	151080	15	AP005726
C 29	29.8	23.1	153065	14	AP005094
C 30	29.6	22.9	436	3	DQ094234
C 31	29.6	22.9	2740	15	AK110305
C 32	29.6	22.9	56276	7	BX276099
C 33	29.6	22.9	110547	5	AF429315
C 34	29.6	22.9	125020	8	AF429315
C 35	29.6	22.9	164179	8	AC116901
C 36	29.6	22.9	174118	8	AC024405
37	29.6	22.9	185341	14	AC069403
38	29.6	22.9	230664	14	AC097977
39	29.4	22.8	870	6	AR480420
40	29.4	22.8	885	6	AR586187
41	29.4	22.8	3520	6	CQ412044
42	29.4	22.8	16853	1	AF246897
C 43	29.4	22.8	37083	8	AC092021
44	29.4	22.8	115469	8	AL356219
45	29.4	22.8	188183	8	AC105921

ALIGNMENTS

RESULT 1	AB072446	600 bp	DNA	linear	BCT 15-JAN-2003
LOCUS	Bifidobacterium longum biovar Longum hup gene for histone-like HU				
DEFINITION	protein HBI, complete cds.				
ACCESSION	AB072446				
VERSION	AB072446.1	GI:19918953			
KEYWORDS	Bifidobacterium longum biovar Longum				
SOURCE	Bifidobacterium longum biovar Longum				
ORGANISM	Bacteria; Actinobacteria; Actinobacteridae; Bifidobacteriales; Bifidobacteriaceae; Bifidobacterium.				
REFERENCE	1	Takeuchi, A., Matsumura, H. and Kano, Y.			
AUTHORS	Cloning and expression in Escherichia coli of a gene, hup, encoding the histone-like protein HU of Bifidobacterium longum				
TITLE	Biosci. Biotechnol. Biochem. 66 (3), 598-603 (2002)				
JOURNAL	12005055				
PUBMED	2 (bases 1 to 600)				
REFERENCE	2 (bases 1 to 600)				
AUTHORS	Kano, Y.				
TITLE	Direct Submission				
JOURNAL	Submitted (03-OCT-2001) Yasunobu Kano, Kyoto Pharmaceutical University, Institute of Molecular and Cellular Biology for Pharmaceutical Sciences; 1, Shichono-cho, Misasagi, Yamashina-ku, Kyoto 6078412, Japan (E-mail: ykano@mb.kyoto-phu.ac.jp, Tel:81755954719, Fax:81755954799)				
FEATURES	Location/Qualifiers				
source	1. .600				
	/organism="Bifidobacterium longum biovar Longum"				
	/mol_type="genomic DNA"				
	/db_xref="taxon:1679"				
RBS	180..185				
gene	193..474				
CDS	/gene="hup"				
	/codon_start=1				
	/transl_table=11				
	/product="histone-like HU protein HBI"				
	/protein_id="BAB88690.1"				
	/db_xref="GI:19918954"				

/translation="MAYKSDLVSKIAOKSNLTKAQAEEAVNAFQDVFVEAMKSGEGL  
KLTLGSPSAERVKRPTGRNPRTEQIDIPASVGRISAGSLKKAITE"

ORIGIN

Query Match 100.0%; Score 129; DB 1; Length 600;  
Best Local Similarity 100.0%; Pred. No. 7.7e-30;  
Matches 129; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
  
QY 1 TGACCTTCTGCTAGCGATTACTTCGAGCATTTACTGACGACAAAGACCCCGACCGAGA 60  
DB 472 TGACCTTCTGCTAGCGATTACTTCGAGCATTTACTGACGACAAAGACCCCGACCGAGA 531  
  
QY 61 TGGTCGGGGCTTTTGTGCGTGTGTCGACGTTGTCCAAACCGATTATTCGGGACT 120  
DB 532 TGGTCGGGGCTTTTGTGCGTGTGTCGACGTTGTCCAAACCGATTATTCGGGACT 591  
  
QY 121 AGTTCAGCG 129  
DB 592 AGTTCAGCG 600

RESULT 2

BD144660 600 bp DNA linear PAT 17-JAN-2003  
LOCUS Drug for gene therapy using anaerobic bacteria.  
DEFINITION  
ACCESSION BD144660  
VERSION BD144660.1 GI:27850418  
KEYWORDS JP 2002097144-A/1.  
SOURCE Bifidobacterium longum  
ORGANISM Bifidobacterium longum  
Bacteria; Actinobacteria; Actinobacteridae; Bifidobacteriales;  
Bifidobacteriaceae; Bifidobacterium.  
REFERENCE 1 (bases 1 to 600)

Fujimori, M., Taniguchi, S., Amano, J., Yazawa, K., Kano, Y.,  
Nakamura, T. and Sasaki, T.  
Drug for gene therapy using anaerobic bacteria  
Patent: JP 2002097144-A 1 02-APR-2002;  
JUN AMANO, MINORU FUJIMORI  
OS Bifidobacterium longum  
PN JP 2002097144-A/1  
PD 02-APR-2002  
PF 21-SEP-2001 JP 2001290187  
PI MINORU FUJIMORI, SHUNICHIRO TANIGUCHI, JUN  
AMANO, KAZUTORA YAZAWA  
PI YASUMASA KANO, TOSHIYUKI NAKAMURA, TAKAYUKI SASAKI PC  
A61K35/74, A61K31/396, A61K31/7068, A61K31/711, A61K38/00, A61K38/00, A61K48/00,  
PC A61P35/00, C12N1/21, C12N15/09// (C12N1/21, C12R1:01), C12N15/00,  
PC A61K37/02,  
PC A61K37/48,  
CC Drug for gene therapy using anaerobic bacteria FH Key  
Location/Qualifiers  
FT CDS (193).. (471).  
Location/Qualifiers  
FEATURES  
source 1..600  
/organism="Bifidobacterium longum"  
/mol\_type="genomic DNA"  
/db\_xref="taxon:216816"

ORIGIN

Query Match 100.0%; Score 129; DB 6; Length 600;  
Best Local Similarity 100.0%; Pred. No. 7.7e-30;  
Matches 129; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
  
QY 1 TGACCTTCTGCTAGCGATTACTTCGAGCATTTACTGACGACAAAGACCCCGACCGAGA 60  
DB 472 TGACCTTCTGCTAGCGATTACTTCGAGCATTTACTGACGACAAAGACCCCGACCGAGA 531  
  
QY 61 TGGTCGGGGCTTTTGTGCGTGTGTCGACGTTGTCCAAACCGATTATTCGGGACT 120  
DB 532 TGGTCGGGGCTTTTGTGCGTGTGTCGACGTTGTCCAAACCGATTATTCGGGACT 591  
  
QY 121 AGTTCAGCG 129

DB 592 AGTTCAGCG 600

RESULT 3  
AE014295.22/c  
WPCOMMENT

Sequence split into 23 fragments LOCUS AE014295 Accession AE014295  
Fragment Name Begin End  
AE014295\_00 1 110000  
AE014295\_01 100001 210000  
AE014295\_02 200001 310000  
AE014295\_03 300001 410000  
AE014295\_04 400001 510000  
AE014295\_05 500001 610000  
AE014295\_06 600001 710000  
AE014295\_07 700001 810000  
AE014295\_08 800001 910000  
AE014295\_09 900001 1010000  
AE014295\_10 1000001 1110000  
AE014295\_11 1100001 1210000  
AE014295\_12 1200001 1310000  
AE014295\_13 1300001 1410000  
AE014295\_14 1400001 1510000  
AE014295\_15 1500001 1610000  
AE014295\_16 1600001 1710000  
AE014295\_17 1700001 1810000  
AE014295\_18 1800001 1910000  
AE014295\_19 1900001 2010000  
AE014295\_20 2000001 2110000  
AE014295\_21 2100001 2210000  
AE014295\_22 2200001 2256640  
Continuation (23 of 23) of AE014295 from base 2200001 (AE014295 Bifidobacterium longum

Query Match 100.0%; Score 129; DB 1; Length 56640;  
Best Local Similarity 100.0%; Pred. No. 7.5e-30;  
Matches 129; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 TGACCTTCTGCTAGCGATTACTTCGAGCATTTACTGACGACAAAGACCCCGACCGAGA 60  
DB 41386 TGACCTTCTGCTAGCGATTACTTCGAGCATTTACTGACGACAAAGACCCCGACCGAGA 41327  
  
QY 61 TGGTCGGGGCTTTTGTGCGTGTGTCGACGTTGTCCAAACCGATTATTCGGGACT 120  
DB 41326 TGGTCGGGGCTTTTGTGCGTGTGTCGACGTTGTCCAAACCGATTATTCGGGACT 41267  
  
QY 121 AGTTCAGCG 129  
DB 41266 AGTTCAGCG 41258

RESULT 4  
AX492788/c  
LOCUS AX492788 156638 bp DNA linear PAT 26-SEP-2002  
DEFINITION Sequence 1106 from Patent EP1227152.  
ACCESSION AX492788  
VERSION AX492788.1 GI:23338471  
KEYWORDS  
SOURCE  
ORGANISM Bifidobacterium longum biovar Longum  
Bifidobacterium longum biovar Longum  
Bacteria; Actinobacteria; Actinobacteridae; Bifidobacteriales;  
Bifidobacteriaceae; Bifidobacterium.

REFERENCE 1  
AUTHORS None.  
TITLE Bacterial strain and genome of bifidobacterium  
JOURNAL Patent: EP 1227152-A 1106 31-JUL-2002;  
Societe des Produits Nestle S.A. (CH)  
FEATURES  
source 1..156638  
/organism="Bifidobacterium longum biovar Longum"  
/mol\_type="unassigned DNA"  
/db\_xref="taxon:1679"  
/note="1106 seq in place of 1099 because seq 1 (2.256.638

bases splitted  
in 7 more sequences.

seq 0001: from 0.000.001 to 0.349.980 length: 349980  
seq 1100: from 0.300.001 to 0.649.980 length: 349980  
seq 1101: from 0.600.001 to 0.949.980 length: 349980  
seq 1102: from 0.900.001 to 1.249.980 length: 349980  
seq 1103: from 1.200.001 to 1.549.980 length: 349980  
seq 1104: from 1.500.001 to 1.849.980 length: 349980  
seq 1105: from 1.800.001 to 2.149.980 length: 349980  
seq 1106: from 2.100.001 to 2.256.638 length: 156638"

## ORIGIN

Query Match 100.0%; Score 129; DB 6; Length 156638;  
Best Local Similarity 100.0%; Pred. No. 7.4e-30;  
Matches 129; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 TGACCTTCTGCTAGCGATTACTTCGACGATTACTGACGACAAAGACCCCGACGAGA 60  
|||||  
Db 141691 TGACCTTCTGCTAGCGATTACTTCGACGATTACTGACGACAAAGACCCCGACGAGA 141632  
|||||  
QY 61 TGGTCGGGGTCTTTTGTGTGTGCTGTGACGCTGTGTCACCGGTATTATTCGGGACT 120  
|||||  
Db 141631 TGGTCGGGGTCTTTTGTGTGTGCTGTGACGCTGTGTCACCGGTATTATTCGGGACT 141572  
|||||

QY 121 AGTTCAGCG 129

Db 141571 AGTTCAGCG 141563

## RESULT 5

AX553955/c  
LOCUS Sequence 9 from Patent WO02074798. 156646 bp DNA linear PAT 27-NOV-2002  
DEFINITION AX553955  
ACCESSION AX553955  
VERSION AX553955.1 GI:25897908  
KEYWORDS Bifidobacterium longum biovar Longum  
SOURCE Bifidobacterium longum biovar Longum  
ORGANISM Bacteria; Actinobacteria; Actinobacteridae; Bifidobacteriales;  
Bifidobacteriaceae; Bifidobacterium.

REFERENCE 1 Arigoni,F., Delley,M., Mollet,B., Pridmore,R.D., Schell,M.A.,  
Pohl,T.G. and Zwhalen,M.C.  
The genome of a bifidobacterium  
Patent: WO 02074798-A 9 26-SEP-2002;  
SOCIETE DES PRODUITS NESTLE S.A. (CH)  
Location/Qualifiers  
1. .156646  
/organism="Bifidobacterium longum biovar Longum"  
/mol\_type="unassigned DNA"  
/db\_xref="taxon:1679"

/notes="seq 1: original length 2256646 splitted into  
seq 1: from 0.000.001 to 0.349.980  
seq 3: from 0.300.001 to 0.649.980  
seq 4: from 0.600.001 to 0.949.980  
seq 5: from 0.900.001 to 1.249.980  
seq 6: from 1.200.001 to 1.549.980  
seq 7: from 1.500.001 to 1.849.980  
seq 8: from 1.800.001 to 2.149.980  
seq 9: from 2.100.001 to 2.256.646"

## ORIGIN

Query Match 100.0%; Score 129; DB 6; Length 156646;  
Best Local Similarity 100.0%; Pred. No. 7.4e-30;  
Matches 129; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 TGACCTTCTGCTAGCGATTACTTCGACGATTACTGACGACAAAGACCCCGACGAGA 60  
|||||  
Db 141699 TGACCTTCTGCTAGCGATTACTTCGACGATTACTGACGACAAAGACCCCGACGAGA 141640  
|||||

QY 61 TGGTCGGGGTCTTTTGTGTGTGCTGTGACGCTGTGTCACCGGTATTATTCGGGACT 120

Db 141639 TGGTCGGGGTCTTTTGTGTGTGCTGTGACGCTGTGTCACCGGTATTATTCGGGACT 141580

QY 121 AGTTCAGCG 129  
|||||  
Db 141579 AGTTCAGCG 141571  
|||||  
RESULT 6  
BX294151/c  
LOCUS Pirellula sp. strain 1 complete genome; segment 19/24. 295650 bp DNA linear BCT 17-APR-2005  
DEFINITION BX294151 BX119912  
ACCESSION BX294151.1 GI:32446812  
VERSION Complete genome.  
KEYWORDS Rhodopirellula baltica SH 1  
ORGANISM Bacteria; Planctomycetes; Planctomycetacia; Planctomycetales;  
Planctomycetaceae; Rhodopirellula.  
1 (bases 1 to 295650)  
Glockner,F.O., Kube,M., Bauer,M., Teeling,H., Lombardot,T.,  
Ludwig,W., Gade,D., Beck,A., Borzym,K., Heitmann,K., Rabus,R.,  
Schlesner,H., Amann,R. and Reinhardt,R.  
Complete genome sequence of the marine planctomycete Pirellula sp.  
strain 1  
Proc. Natl. Acad. Sci. U.S.A. 100 (14), 8298-8303 (2003)  
12835416  
2 (bases 1 to 295650)  
Kube,M., Borzym,K., Heitmann,K., Klages,S., Marquardt,I.,  
Lehrack,S., Beck,A., Pawlik,R., Reinhardt,R., Gloeckner,F.O.,  
Bauer,M., Teeling,H., Lombardot,T., Ludwig,W., Gade,D., Rabus,R.,  
Schlesner,H. and Amann,R.  
Direct Submission  
Submitted (21-JAN-2003) Max Planck Institute for Molecular  
Genetics, proScience Ihnestrasse 73, D-14195 Berlin, Germany Max  
Planck Institute for Marine Microbiology Celsiusstrasse 1, D-28359  
Bremen, Germany  
This project was carried out by  
\*Max Planck Institute for Molecular Genetics, Berlin, Germany; \*Max  
Planck Institute for Marine Microbiology, Bremen, Germany; in the  
framework of the REGX-project, <http://www.regx.de> -----  
Genome Center  
Center: Max Planck Institute for Molecular Genetics  
----- Summary Statistics  
Sequencing vector: pUC19; 100% of reads  
Chemistry: Dye-terminator Big Dye; 100% of reads  
Assembly program: Phrap; version 0.990329  
Consensus quality: 7142841 bases at least Q40  
Consensus quality: 7145138 bases at least Q30  
Consensus quality: 7145484 bases at least Q20  
Quality coverage: 8.03  
-----  
This sequence was finished as follows unless otherwise noted: all  
regions were double stranded, sequenced with an alternate  
chemistry, or covered by high quality data (i.e., phred quality >=  
30); an attempt was made to resolve all sequencing problems, such  
as compressions and repeats; all regions were covered by at least  
one plasmid sequence; assembly was additionally confirmed by long  
range pcr and cosmid end sequences.  
-----  
See <http://www.micro-genomes.mpg.de/pirellula/> for more information  
including minimal tiling path from a set of 220 cosmids out of  
908. See the misc feature tag below for the boundaries of the MTP  
cosmids. ----- Annotation  
Center: Max Planck Institute for Marine Microbiology  
Celsiusstrasse 1, D-28359 Bremen, Germany.  
Center Code: MPIMM  
Email: [foe@mpi-bremen.de](mailto:foe@mpi-bremen.de)  
Phone: +49 (0)421 2028 938 Fax: +49 (0)421 2028 580  
-----  
Three different programs (Glimmer, Critica and Orpheus) were used  
for ORF-prediction. A nonredundant list of ORFs was generated by  
suitable parsing of the results.  
Automated annotation was done with the software package Pedant Pro

(http://www.biomax.de). All ORF predictions and annotations were manually corrected by considering all results of the different tools applied. See http://www.regx.de for more information and access to supplementary information.

FEATURES	Location/Qualifiers	
	source	
gene	1..295650	/organism="Rhodopirellula baltica SH 1"
		/mol_type="genomic DNA"
		/strain="1"
		/db_xref="taxon:243090"
		/complement(414..1508)
CDS		/genes="pknH"
		/locus_tag="RB10133"
		/complement(414..1508)
		/gene="pknH"
		/locus_tag="RB10133"
gene		/EC number="2.7.1.-"
		/functions="unspecified signal transduction; unspecified kinase or ATP dependent regulatory protein; cellular communication/signal transduction"
		/note="best DB hits: BLAST: swissprot:Q11053; PKNH MYCTU PROBABLE SERINETHREONINE-PROTEIN; E=3e-35 emb1: CAB94054.1; (AL358672) putative serinethreonine-protein; E=1e-34 emb1:CAA10713.1; (AJ132604) hypothetical protein [Lactococcus; E=2e-33 COG: RV1266c; COG0515 serine/threonine protein kinases; E=3e-36 PFAM: PF00069; Protein kinase domain; E=7.6e-54"
		/codon_start=1
		/transl_table=1
gene		/product="probable serine/threonine-protein kinase pknH"
		/protein_id="CAD78719.1"
		/db_xref="GI:32446813"
		/db_xref="GOA:Q7UFG3"
		/db_xref="InterPro:IPR000719"
CDS		/db_xref="InterPro:IPR008271"
		/db_xref="UniProt/TREMBL:Q7UFG3"
		/translation="MPHLRLTDFELGAVLGVTGYDGKILDDVEVHPAAEAIRG GDAVKKLPAPVQDDLIQARFRREMILRQHPNIGYFGGSDGQIFYNVERVD QGTIKOLLETGALANPVVDVARQVCSALQCAHGHVHRDKPGLFLTRDAHVKL DFGFIARDHSSDLSQGLTVGTTHAYMAPEQITGDEAISGKADLYALGCVLFEMLANR KVFGEANFAQLFQHLRTKAPTASIVPDVPPELSQVIADECLSKSPDRPFNARSVQG VMIRIGKYNLSPDPSVGVNHSCTGNAPKDVSAHSVTEKRRLLLEEQIHYRLGSGTS REPVSAGKVSILVAILVILVAVLSGGG"
		/complement(1529..1843)
		/locus_tag="RB10134"
gene		/locus_tag="RB10134"
		/functions="ribosomal proteins"
		/note="best DB hits: BLAST: gb:AAK05177.1; AE006340.10 (AE006340) 50S ribosomal protein L21; E=4e-10 pir:F82920; ribosomal protein L21 U0212 [imported] - Ureaplasma; E=3e-11 gb:AAK35614.1; (AF041468) ribosomal protein L21 [Guillardia; E=3e-10 COG: RV2442c; COG0261 Ribosomal protein L21; E=3e-07 PFAM: PF00829; Ribosomal prokaryotic L21 prot; E=2.2e-36"
		/codon_start=1
		/transl_table=1
CDS		/product="50S ribosomal protein L21"
		/protein_id="CAD78720.1"
		/db_xref="GI:32446814"
		/db_xref="GOA:Q7UFG2"
		/db_xref="InterPro:IPR001787"
gene		/db_xref="UniProt/TREMBL:Q7UFG2"
		/translation="MYAIFVDGGRQYRVEFGMELVDYRDIAAGENLKFTVLAVGAE DGLKLGAPTLDGVSASVLGMSQDKIYIKFRRRKHSKKRTGHRKQNRIRIEIA GV"
		/complement(1767..2018)
		/locus_tag="RB10135"
		/complement(1767..2018)
CDS		/locus_tag="RB10135"
		/codon_start=1
		/transl_table=1

gene		/product="hypothetical protein"
		/protein_id="CAD78721.1"
		/db_xref="GI:32446815"
		/db_xref="UniProt/TREMBL:Q7UFG1"
		/translation="MLDRPIFGSTDLFTSDDLLSLAKNQWPKRRSRVVILRDSKIR PSQQFPLHPPEFSPRHVCHLCRRWTFIPSRARNGTRRRLP"
CDS		/complement(2151..2372)
		/locus_tag="RB10136"
		/complement(2151..2372)
		/locus_tag="RB10136"
		/codon_start=1
gene		/transl_table=1
		/product="hypothetical protein"
		/protein_id="CAD78722.1"
		/db_xref="GI:32446816"
		/db_xref="UniProt/TREMBL:Q7UFG0"
CDS		/translation="MLVLRNRSELSVGSGAARAARCVTGGIAPNGTYFTPPGGGCEP LRASPVTP"
		/complement(2151..2372)
		/locus_tag="RB10137"
		/complement(2151..2372)
		/locus_tag="RB10137"
gene		/codon_start=1
		/transl_table=1
		/product="hypothetical protein-putative conserved hypothetical protein"
		/protein_id="CAD78723.1"
		/db_xref="GI:32446817"
CDS		/db_xref="InterPro:IPR011481"
		/db_xref="UniProt/TREMBL:Q7UFG9"
		/translation="MSRSRFGVLYDAPLKRSLGGGRATPPFRRTRGRGSPMENGADC PPEITSLNARPTPPAARAGFSRRYWRCT"
		/complement(2478..3281)
		/locus_tag="RB10138"
gene		/complement(2478..3281)
		/locus_tag="RB10138"
		/function="inner membrane organization"
		/note="best DB hits: BLAST: Pir.T34929; hypothetical protein SC3P9.07 SC3P9.07 - Streptomyces; E=0.018"
CDS		/codon_start=1
		/transl_table=1
		/product="conserved hypothetical protein-putative membrane lipoprotein"
		/protein_id="CAD78724.1"
		/db_xref="GI:32446818"
gene		/db_xref="UniProt/TREMBL:Q7UFG8"
		/translation="MRLVTDFFNAFPFLQRNSTSSALMSLVFAGCCLLVNPFPSIAEA DPAEQTSNVSVKIFDTNAPGWRQLDGDFFRVNGDDQTLTWEGTALGSGQPIGVTR TNPEVKNFELVQWKHLKPAGNSGVPAWPMASALKDLPPNRLPNTGTIEVQMLDLYGR MYTEKTKGPTWFTSHGDIIPAVGKSSMQPPPLSPDGHRSFPSPAETNPHGWNQYVY RGINREIRLVNNGVEVSGRSQSPDGFCLSESGSPIRFREIWLREL"
		/complement(3382..3621)
		/locus_tag="RB10139"
		/complement(3382..3621)
CDS		/locus_tag="RB10139"
		/codon_start=1
		/transl_table=1
		/product="hypothetical protein"
		/protein_id="CAD78725.1"
gene		/db_xref="GI:32446819"
		/db_xref="UniProt/TREMBL:Q7UFG7"
		/translation="MRKHDRMKIRGISKNPIDLPSRYRGRVLKPKKRRSRFSFLRE
Query Match 25.3%; Score 32.6; DB 1; Length 295650;		
Best Local Similarity 53.5%; Pred. No. 24;		
Matches 68; Conservative 0; Mismatches 59; Indels 0; Gaps 0;		
Qy	1	TGACCTTCTCGTAGCGATTACTTCGAGCATTACTGACGACAAAGACCCCGACCGAGA 60
Db	148951	TGACGTAGCCTCGATCGAATAGTTCGAACGAGGACGACGACCAACGTCAGA 148892
Qy	61	TGGTCGGGGTCTTTTGTGCTGCTGACGTGTGTCACCGCTATTATTCGGACT 120

Db 148891 TGTGTGCTGATTGCGGAGTTTCGGATGCGAGTCCGAAAGAGCATTCCTCTACCGAACT 148832

QY 121 AGTTCAG 127

Db 148831 GTTTCGG 148825

RESULT 7  
CNS07EGA/c  
LOCUS  
DEFINITION  
CNS07EGA 194439 bp DNA linear INV 16-APR-2005  
Chromosome III of strain GB-M1 of Encephalitozoon cuniculi  
(Microspora).

ACCESSION  
AL590443 GI:13560062

VERSION  
AL590443.1 GI:13560062

KEYWORDS  
AIDS opportunistic pathogen; Genome sequence; intracellular  
parasite.

SOURCE  
Encephalitozoon cuniculi GB-M1

ORGANISM  
Encephalitozoon cuniculi GB-M1

REFERENCE  
1 (bases 1 to 194439)

AUTHORS  
Katinka M.D., Duprat S., Cornillot E., Metenier G., Thomarat P.,  
Prensier G., Barbe V., Peyretailade E., Brottier P., Wincker P.,  
Delbac F., El Alaoui H., Peyret P., Saurin W., Gouy M.,  
Weissenbach J. and Vivares C.P.

TITLE  
Encephalitozoon cuniculi

JOURNAL  
Nature 414 (6862), 450-453 (2001)

PUBMED  
11719806

REFERENCE  
2 (bases 1 to 194439)

AUTHORS  
Direct Submission

TITLE  
Submitted (05-APR-2001) Genoscope - Centre National de Sequencage,  
2 rue Gaston Cremieux, CP 5706, 91057 EVRY cedex, FRANCE. (E-mail :  
seqref@genoscope.cns.fr - Web : www.genoscope.cns.fr)

COMMENT  
Corresponding author : Equipe de Parasitologie Moleculaire et  
Cellulaire, UMR CNRS 6023, Universite Blaise Pascal, 63177 Aubiere  
cedex, France E-mail : Christian.VIVARES@bpcclermont.fr

Web : http://www.protistes.univ-bpclermont.fr/fr/parasito.htm The  
submitted sequence encompasses the coding core of the chromosome.  
The telomeric and subtelomeric sequences are not included in this  
submission.

FEATURES  
Location/Qualifiers

1. .194439

/organism="Encephalitozoon cuniculi GB-M1"

/mol\_type="genomic DNA"

/strain="GB-M1"

/db\_xref="taxon:284813"

/chromosome="III"

550..1380

/gene="ECU03\_0010"

/genes="ECU03\_0010"

/function="unknown"

/note="03\_0010, unknown, gene found by Glimmer [Bad Olap 2

167 0]"

/codon\_start=1

/product="hypothetical protein"

/protein\_id="CAD26148.1"

/db\_xref="GI:19168629"

/db\_xref="UniProt/Swiss-Prot:Q8SW85"

/translation="MAAQTLPETAKPKHNSQSEAGPAPLASPTAMPRPASHLADMP  
SDHPDRSKARCPPTNCGTFKQPPSVAATSRPKGPNFLQPTKGTPTPKK  
KNHTGCTHREANPENYKHTETEPPIISCHPPPGTATPNLLPCPNTPSSFCQ  
TRDSPLPORNMWHPETPLTHRHSPSQIPRIPIPPYNLSTAKTYLQPPYLPPI  
LSHEITHFVLLCPYEHHTNRAAREPATLENNPRHRIRLNIYIPSHVPHLRQF  
"

1211..2020

/gene="ECU03\_0020"

1211..2020

/genes="ECU03\_0020"

/function="unknown"

/note="03\_0020, unknown, gene found by Glimmer [Shorter 1

167 0]"

/codon\_start=1

/product="hypothetical protein"

/protein\_id="CAD26149.1"

/db\_xref="GI:19168630"

/db\_xref="UniProt/Swiss-Prot:Q8SW84"

/translation="MKSISYLSFAHMTHTTEQHAENOPHMTKILDIAPFVSITFP  
AMCLIFDSDSEESPFRLFTILLPFSYSAVOYALLYTNMKNHNPETILHTLYT  
LSLLIAPFTIISLIIIPSLNEMWHAASFFPIVLPSFTVPAYLLSSYFLVPRQI  
RLTDVISLISVCSIVNLLVFKENYYPYSALISSISVLLQLLSEKCHLQKQSPPS  
TASSRAVLLIHLAVLYTELGYCAIYLLDDHFLHLLKMKSLPSEPHQ"

complement(3222..3578)

/gene="ECU03\_0030"

complement(3222..3578)

/gene="ECU03\_0030"

/function="unknown"

/note="03\_0030, unknown, gene found by Glimmer"

/codon\_start=1

/product="hypothetical protein"

/protein\_id="CAD26150.1"

/db\_xref="GI:19168631"

/db\_xref="UniProt/Swiss-Prot:Q8STE8"

/translation="MCFPVGGJLPYLALRSVAYICVFGSSRAPDLIRAFESPHFKR  
FDGAGTYKAKGRCGRFADVVNGVAGQISDMADKVGGEPEVYCIWKKRGEVEMLLK  
VKEYRKGYSKGRRR"

4169..4975

/gene="ECU03\_0040"

4169..4975

/gene="ECU03\_0040"

/function="unknown"

/note="03\_0040, unknown, gene found by Glimmer"

/codon\_start=1

/product="hypothetical protein"

/protein\_id="CAD26151.1"

/db\_xref="GI:19168632"

/db\_xref="InterPro:IPR011004"

/db\_xref="UniProt/Swiss-Prot:Q8SW83"

/translation="MNTTVPSPHRTQCHTENLQHWKILGIAPVISAPPAIMYFIF  
TKDSDPSDLRFITILLSPFSYSAVOYALLYTNMKNHNPETILHTLYTNLLLL  
AFSIIISLITTLPIKWNKDGPILSIFLPLMSPAYLLSTCRLVPGQIGTDT  
GNVLIDITLCSVGSLLILDESSYCYFAIISILIRLLGSLKLSPEKSPSPPT  
APWRIAVFVLIIFAAIYAFMMVMSIDILSDHFDLLTKARSTSVSKPGQ"

5400..6665

/gene="ECU03\_0050"

5400..6665

/gene="ECU03\_0050"

/function="unknown"

/note="03\_0050, unknown, gene found by Glimmer"

/codon\_start=1

/product="hypothetical protein"

/protein\_id="CAD26152.1"

/db\_xref="GI:19168633"

/db\_xref="InterPro:IPR001664"

/db\_xref="UniProt/TREMBL:Q8SW82"

/translation="MEQISRIKTLTDEVDLTDYDERAFRLGSVEKLVAVRAFWLISDIVK  
CKMHPATACRYNLLDSILEDEEKHPEGITAVNAKDARLALBDVNRKCOILSLS  
KXAGTEPPYFGAIDGDKYGLLKAFKDALRKVLPHVGRDLKVVNMKLEAAMS  
DFELVKQLDGEKQVERPSTLDIYKLMNEVNEVEVPLKSVIRDLGLGEGNYNFEAN  
WSFYSPYRDINEAALYIFOKYFQHDIDEASLRTIKNELECEKVIKVPKSWRLQD  
RLRDVRTAFDSLSQKLKALRTSCTRIIDVDVESNGKSPDSFSELNGLMREAKGLSD  
LKERSTIDLERVLPSPPMKLCFLCFMFIALLIHSFLHAEKNSRVRTKRWNTALYS  
LAVAGTIDHQMVLVGGGTGMRMGQVMC"

6782..7153

/gene="ECU03\_0060"

6782..7153

/gene="ECU03\_0060"

/function="unknown"

/note="03\_0060, unknown, gene found by Glimmer [Delay by

228 5 109 38]"

/codon\_start=1

/product="hypothetical protein"

/protein\_id="CAD26153.1"





rice.

URL : <http://cdna01.dna.affrc.go.jp/cDNA/>  
 NIAS Rice Full-Length cDNA Project Team: Kikuchi, S., Satoh, K., Nagata, T., Kawagashira, N., Doi, K., Kishimoto, N., Yazaki, J., Ishikawa, M., Yamada, H., Ooka, H., Hotta, I., Kojima, K., Namiki, T., Ohneda, E., Yahagi, W., Suzuki, K., Li, C., Ohtsuki, K., Shishiki, T., Yamamoto, M. and Nakahama, Y.  
 FAIS Genome Sequencing & Analysis Group: Ootomo, Y., Iida, Y., Fujimura, T., Ikeda, R., Ishibiki, J., Kawamata, M., Kobayashi, M., Kodama, T., Kurosaki, T., Kusumegi, T., Lu, M., Masuda, H., Miura, J., Mizuno, K., Narikawa, R., Niihara, J., Oka, M., Ryu, R., Sugano, S., Sugiyama, A., Suzuki, Y., Teunoda, Y., Ueda, M., Xie, Q., Yokomizo, S., Yoshimura, A., Matsubara, K. and Murakami, K.  
 Genome Exploration Research Group in Riken Genomic Sciences Center and Genome Science Laboratory in Riken: Adachi, J., Aizawa, K., Akimura, T., Arakawa, T., Carninci, P., Fukuda, S., Hanagaki, T., Hara, A., Hashizume, W., Hayaashida, K., Hayatsu, N., Hiramoto, K., Hiraoka, T., Hori, F., Iida, J., Imamura, K., Imotani, K., Ishii, Y., Itoh, M., Kagawa, I., Kangawa, S., Katoh, H., Kawai, J., Kishikawa-Hirozane, T., Kojima, Y., Kondo, S., Konno, H., Kouda, M., Koya, S., Kurihara, C., Matsuyama, T., Miyazaki, A., Murata, M., Nakamura, M., Nishi, K., Nomura, K., Numasaki, R., Ohno, M., Osato, N., Ota, Y., Saitoh, H., Sakai, C., Sakai, K., Sakazume, N., Sano, H., Sasaki, D., Sato, K., Shibata, K., Shinagawa, A., Shiraki, T., Sogabe, Y., Tagami, M., Tagami-Takeda, Y., Tagawa, A., Takahashi, F., Takaku-Akahira, S., Tanaka, T., Tomaru, A., Toya, T., Waki, K., Yasunishi, A. and Hayashizaki, Y.

FEATURES  
 source 1. 3048  
 Location/Qualifiers  
 /organism="Oryza sativa (japonica cultivar-group)"  
 /mol\_type="mRNA"  
 /cultivar="Nipponbare"  
 /db\_xref="taxon:39947"  
 /clone="002-178-D02"

## ORIGIN

Query Match 24.7%; Score 31.8; DB 15; Length 3048;  
 Best Local Similarity 61.4%; Pred. No. 44;  
 Matches 51; Conservative 0; Mismatches 32; Indels 0; Gaps 0;  
 QY 16 AGCGATTCTCGACGATTTACTGACGACAAAGACCCGACGAGGTGCGGGTCTTTT 75  
 Db 506 AGTTATCACAGCATGGCTCTCTCACACCTGACCGCCACGAGCTGCTTCAGCTGTT 565  
 QY 76 TGTGTGGTGTGTGACGTGTTG 98  
 Db 566 TGTGTGGTGTGTGTTGTTT 588

RESULT 9  
 AC138542/c 90417 bp DNA linear HTG 20-MAR-2004  
 LOCUS Magnaporthe grisea chromosome 7 clone 12G19, \*\*\* SEQUENCING IN  
 DEFINITION PROGRESS \*\*\*

ACCESSION AC138542  
 VERSION AC138542.2 GI:45597487  
 KEYWORDS HTG; HTGS PHASE2.  
 SOURCE Magnaporthe grisea (anamorph: Pyricularia grisea)  
 ORGANISM Magnaporthe grisea

REFERENCE 1 (bases 1 to 90417)  
 AUTHORS Eukaryota; Fungi; Ascomycota; Pezizomycotina; Sordariomycetes; Sordariomycetes incertae sedis; Magnaportheaceae; Magnaporthe.  
 REFERENCE 2 (bases 1 to 90417)  
 AUTHORS Thon, M.R., Mitchell, T., Brown, D.E., Diener, S., Taro, A., Pan, H. and Dean, R.A.

TITLE The sequence of Magnaporthe grisea chromosome 7  
 JOURNAL Unpublished  
 REFERENCE 2 (bases 1 to 90417)  
 JOURNAL Dean, R.A. Dr. Mitchell, T. Dr. Thon, M. Dr and Brown, D.E.  
 AUTHORS Direct Submission  
 TITLE Submitted (09-JAN-2003) Plant Pathology - Fungal Genomics Laboratory, North Carolina State University, 840 Main Campus Drive, Raleigh, NC 27606, USA

## REFERENCE

## AUTHORS

## TITLE

## JOURNAL

## COMMENT

3 (bases 1 to 90417)  
 Thon, M.R., Mitchell, T., Brown, D.E., Diener, S., Taro, A., Pan, H. and Dean, R.A.  
 Direct Submission  
 Submitted (20-MAR-2004) Center for Integrated Fungal Research, North Carolina State University, Plant Pathology - 840 Main Campus Dr, Raleigh, NC 27606, USA  
 COMMENT On Mar 20, 2004 this sequence version replaced gi:27552536.  
 \* NOTE: This is a 'working draft' sequence. It currently  
 \* consists of 1 contigs. Gaps between the contigs  
 \* are represented as runs of N. The order of the pieces  
 \* is believed to be correct as given, however the sizes  
 \* of the gaps between them are based on estimates that have  
 \* provided by the submitter.  
 \* This sequence will be replaced  
 \* by the finished sequence as soon as it is available and  
 \* the accession number will be preserved.

## FEATURES

source 1. 90417  
 Location/Qualifiers  
 /organism="Magnaporthe grisea"  
 /mol\_type="genomic DNA"  
 /db\_xref="taxon:148305"  
 /chromosomes="7"  
 /clone="12G19"

## ORIGIN

Query Match 24.3%; Score 31.4; DB 14; Length 90417;  
 Best Local Similarity 54.9%; Pred. No. 58;  
 Matches 62; Conservative 0; Mismatches 51; Indels 0; Gaps 0;  
 QY 5 CTTCTCTCGTAGCGATTACTTCGACGATTACTGACGACAAAGACCCGACGAGTGGT 64  
 Db 88659 CTTCCGCTCTCAGGCTCCATCGAACCATCTCCATCCAGGCTCTCGAAGGAGTGGT 88600  
 QY 65 CGGGGTCTTTTGTGTGCTGCTGACGCTGTGTCCACCGTATTATTCGG 117  
 Db 88599 CTTGGCGTCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTG 88547

## RESULT 10

AC138544/c 92770 bp DNA linear HTG 20-MAR-2004  
 LOCUS Magnaporthe grisea chromosome 7 clone 15G13, \*\*\* SEQUENCING IN  
 DEFINITION PROGRESS \*\*\*

## ACCESSION

AC138544 2 GI:45597492  
 VERSION HTG; HTGS PHASE2.  
 KEYWORDS Magnaporthe grisea  
 SOURCE Magnaporthe grisea

## ORGANISM

REFERENCE 1 (bases 1 to 92770)  
 AUTHORS Eukaryota; Fungi; Ascomycota; Pezizomycotina; Sordariomycetes; Sordariomycetes incertae sedis; Magnaportheaceae; Magnaporthe.  
 REFERENCE 2 (bases 1 to 92770)  
 AUTHORS Thon, M.R., Mitchell, T., Brown, D.E., Diener, S., Taro, A., Pan, H. and Dean, R.A.

## TITLE

The sequence of Magnaporthe grisea chromosome 7  
 JOURNAL Unpublished  
 REFERENCE 2 (bases 1 to 92770)  
 JOURNAL Dean, R.A. Dr. Mitchell, T. Dr. Thon, M. Dr and Brown, D.E.

## AUTHORS

Direct Submission  
 Submitted (09-JAN-2003) Plant Pathology - Fungal Genomics Laboratory, North Carolina State University, 840 Main Campus Drive, Raleigh, NC 27606, USA

## REFERENCE

3 (bases 1 to 92770)  
 Thon, M.R., Mitchell, T., Brown, D.E., Diener, S., Taro, A., Pan, H. and Dean, R.A.  
 Direct Submission  
 Submitted (20-MAR-2004) Center for Integrated Fungal Research, North Carolina State University, Plant Pathology - 840 Main Campus Dr, Raleigh, NC 27606, USA  
 COMMENT On Mar 20, 2004 this sequence version replaced gi:27552538.  
 \* NOTE: This is a 'working draft' sequence. It currently  
 \* consists of 1 contigs. Gaps between the contigs

```

* are represented as runs of N. The order of the pieces
* is believed to be correct as given, however the sizes
* of the gaps between them are based on estimates that have
* been provided by the submittor.
* This sequence will be replaced
* by the finished sequence as soon as it is available and
* the accession number will be preserved.
*
* 1 92770: contig of 92770 bp in length.
  Location/Qualifiers
    1..92770
      /organism="Magnaporthe grisea"
      /mol_type="genomic DNA"
      /db_xref="taxon:148305"
      /chromosome="7"
      /clone="15G13"

ORIGIN
Query Match      24.3%; Score 31.4; DB 14; Length 92770;
Best Local Similarity 54.9%; Pred. No. 58;
Matches 62; Conservative 0; Mismatches 51; Indels 0; Gaps 0;

QY 5 CTTCTGCTCAGCATTACTTCGACGATTCGACGAAAGACCCGACCGAGATGGT 64
    |||||
Db 468 CTTCCGCTCTCAGGCTCCATCGAACCATCTCCATACCAGGTCTCTCGAAGGATCGTT 409
    |||||

QY 65 CGGGGCTTTTCTTGTGGTCTGTGACGTTGTCCAACGTTATTATTCGG 117
    |||||
Db 408 CTTGGGGCTTCTGTCTGCTCTCTGTGGTGGCAATCCATCGCGGAAGTCTCTG 356
    |||||

RESULT 11
AC144889
LOCUS AC144889 128428 bp DNA linear HTG 11-JUN-2003
DEFINITION Bos taurus clone RP42-541E10, WORKING DRAFT SEQUENCE, 6 ordered
pieces.
ACCESSION AC144889.2 GI:31581601
VERSION HTG; HTGS_PHASE2; HTGS_DRAFT.
KEYWORDS Bos taurus (cow)
SOURCE
  ORGANISM
    Bos taurus
      Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
      Mammalia; Eutheria; Laurasiatheria; Cetartiodactyla; Ruminantia;
      Pecora; Bovidae; Bovinae; Bos.
REFERENCE
  1 (bases 1 to 128428)
    Antonellis,A., Ayala,K., Beckstrom-Sternberg,S.M., Benjamin,B.,
    Blakeley,R.W., Bouffard,G.G., Brinkley,C., Brooks,S., Cariaga,K.,
    Chu,G., Coleman,B., Coleman,H., Engle,J., Granite,S., Guan,X.,
    Gupta,J., Haghighi,P., Han,J., Hansen,N., Ho,S.-L., Hu,P.,
    Hurle,B., Idol,J.R., Karlins,E., Kwong,P., Laric,P., Lee-Lin,S.-Q.,
    Legaapi,R., Maduro,Q.L., Maduro,V.B., Margulies,E.H., Masello,C.,
    Maskeri,B., McDowell,J., Paguirigan,C., Pearson,R., Portnoy,M.E.,
    Prasad,A., Reddix-Dugue,N., Schandler,K., Schueler,M.G., Shah,K.,
    Sison,C., Stantripo,S., Thomas,J.W., Thomas,P.J., Tsipouri,V.,
    Vogt,J.L., Wetherby,K.D., Wiggins,L., Young,A. and Green,E.D.
    NISC Comparative Sequencing Initiative
  Unpublished
  2 (bases 1 to 128428)
    Green,E.D.
  Direct Submission
  Submitted (23-MAY-2003) NIH Intramural Sequencing Center, 8717
  Government Circle, Gaithersburg, MD 20877, USA
  3 (bases 1 to 128428)
    Green,E.D.
  Direct Submission
  Submitted (11-JUN-2003) NIH Intramural Sequencing Center, 8717
  Government Circle, Gaithersburg, MD 20877, USA
  On Jun 11, 2003 this sequence version replaced gi:31044309.
  ----- Genome Center
    Center: NIH Intramural Sequencing Center
    Center code: NISC
    Web site: http://www.nisc.nih.gov
    Contact: nisc.zoo@nih.gov
  ----- Project Information

```

Center project name: cey  
Center clone name: 541E10

The sequence data in this record represents an 'enhanced' version of a Phase 2 submission. Specifically, the indicated order and orientation of each sequence contig, has been established using one or more of the following: read-pair data from individual subclones, overlaps with neighboring clones, alignment with available reference sequence (e.g., human), and/or confirmation by PCR testing. In addition, the sequence assembly is based on at least 8x average coverage in Q20 bases and has been reviewed to rule out gross misassemblies, the low-quality ends of sequence contigs have been trimmed away, and each base is associated with a Phrap-derived quality score.

#### ----- Summary Statistics

Sequencing vector: plasmid; n/a; 100% of reads  
Chemistry: Dye-terminator Big Dye; 100% of reads  
Assembly program: Phrap; version 0.990319  
Consensus quality: 127382 bases at least Q40  
Consensus quality: 127698 bases at least Q30  
Consensus quality: 127848 bases at least Q20  
Insert size: 130000; agarose-fp  
Insert size: 127928; sum-of-contigs  
Quality coverage: 10.65x in Q20 bases; agarose-fp  
Quality coverage: 10.82x in Q20 bases; sum-of-contigs

-----

\* NOTE: This is a 'working draft' sequence. It currently consists of 6 contigs. Gaps between the contigs are represented as runs of N. The order of the pieces is believed to be correct as given, however the sizes of the gaps between them are based on estimates that have been provided by the submittor.

\* This sequence will be replaced

\* by the finished sequence as soon as it is available and

\* the accession number will be preserved.

\* 1 33969: contig of 33969 bp in length

\* 33970 34069: gap of unknown length

\* 34070 53360: contig of 19291 bp in length

\* 53361 53460: gap of unknown length

\* 53461 63639: contig of 10079 bp in length

\* 63640 112600: contig of 48961 bp in length

\* 112601 112700: gap of unknown length

\* 112701 118744: contig of 6044 bp in length

\* 118745 118845: gap of unknown length

\* 118845 128428: contig of 9584 bp in length.

#### FEATURES

Source

1..128428  
/organism="Bos taurus"  
/mol\_type="genomic DNA"  
/db\_xref="taxon:9913"  
/clone="RP42-541E10"  
/clone\_lib="RP42"

misc\_feature 1..48703  
/note="clone overlaps with GenBank Accession Number AC095020 clone RP42-243E1 (center project name cje)"

misc\_feature 1..33969  
/note="assembly\_fragment  
clone\_end:T7  
vector\_side:left"

gap 33970..34069  
/estimated\_length=unknown

misc\_feature 34070..53360  
/note="assembly\_fragment"

gap 53361..53460  
/estimated\_length=unknown

misc\_feature 53461..63639  
/note="assembly\_fragment"

gap 63640..63639  
/estimated\_length=unknown

misc\_feature 63640..112600  
/note="assembly\_fragment"

```
misc_feature 72761..128428
/notes="clone overlaps with GenBank Accession Number
AC093397 clone RP42-405D15 (center project name cja)"
gap 112601..112700
/estimated length=unknown
misc_feature 112701..118744
/notes="assembly_fragment"
gap 118745..118844
/estimated length=unknown
misc_feature 118845..128428
/notes="assembly_fragment
clone end:SP6
vector_side:right"
ORIGIN
Query Match 24.3%; Score 31.4; DB 14; Length 128428;
Best Local Similarity 53.7%; Pred. No. 58;
Matches 65; Conservative 0; Mismatches 56; Indels 0; Gaps 0;
QY 8 CTGCTGATGAGTACTTTCGAGCATTCGACGACAAAGACCCGACGAGATGTCGG 67
Db 111483 CTCCCGTTTTCCTCCCTCGGCCCTCAGCTCCGACGAGCCTCCACCTGGCTCCCA 111542
QY 68 GGTCTTTTCTGCTGGCTGACGCTGTCACACGTAATTCGCGACTAGTTCAG 127
Db 111543 GGCCTTGTGTTGTAGTTGCTAAGTCATGTCACCTCTGACCCCGGAGCTTCAG 111602
QY 128 C 128
Db 111603 C 111603
RESULT 12
AC093397
LOCUS AC093397.2 GI:21465397
DEFINITION Bos taurus clone RP42-405D15, WORKING DRAFT SEQUENCE, 10 ordered
pieces.
ACCESSION AC093397
VERSION AC093397.2
KEYWORDS HTG; HTGS PHASE2; HTGS_DRAFT.
SOURCE Bos taurus (cow)
ORGANISM Bos taurus
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Laurasiatheria; Cetartiodactyla; Ruminantia;
Fecora; Bovidae; Bovinae; Bos.
REFERENCE 1 (bases 1 to 207821)
AUTHORS Akhter,N., Antonellis,A., Ayele,K., Beckstrom-Sternberg,S.M.,
Benjamin,B., Blakesley,R.W., Bouffard,G.G., Breen,K., Brinkley,C.,
Brooks,S., Dietrich,N.L., Granite,S., Guan,X., Gupta,J.,
Haghighi,P., Hansen,N., Ho,S.-L., Idol,J.R., Karlins,E., Laric,P.,
Lee-Lin,S.-Q., Legaspi,R., Maduro,Q.L., Maduro,V.B.,
Margulies,E.H., Masiello,C., Maskeri,B., Mastrian,S.D.,
McCloskey,J.C., McDowell,J., Paguirigan,C., Pearson,R.,
Portnoy,M.E., Prasad,A., Schueler,M.G., Scantripop,S., Thomas,J.W.,
Wetherby,K.D., Wiggins,L., Tsugeon,C., Vogt,J.L., Walker,M.A.,
Zhang,L.-H. and Green,E.D.
NISC Comparative Sequencing Initiative
Unpublished
REFERENCE 2 (bases 1 to 207821)
AUTHORS Green,E.D.
TITLE Direct Submission
JOURNAL Submitted (22-AUG-2001) NIH Intramural Sequencing Center, 8717
Government Circle, Gaithersburg, MD 20877, USA
REFERENCE 3 (bases 1 to 207821)
AUTHORS Green,E.D.
TITLE Direct Submission
JOURNAL Submitted (19-NOV-2002) NIH Intramural Sequencing Center, 8717
Government Circle, Gaithersburg, MD 20877, USA
COMMENT On Jun 19, 2002 this sequence version replaced gi:15277346.
----- Genome Center
Center: NIH Intramural Sequencing Center
Center code: NISC
Web site: http://www.nisc.nih.gov
```

Contact: nisc\_zoo@nhgri.nih.gov  
----- Project Information  
Center project name: cja  
Center clone name: 405D15

The sequence data in this record represents an 'enhanced' version of a Phase 2 submission. Specifically, the indicated order and orientation of each sequence contig has been established using one or more of the following: read-pair data from individual subclones, overlaps with neighboring clones, alignment with available reference sequence (e.g., human), and/or confirmation by PCR testing. In addition, the sequence assembly is based on at least 8x average coverage in Q20 bases and has been reviewed to rule out gross misassemblies, the low-quality ends of sequence contigs have been trimmed away, and each base is associated with a Phrap-derived quality score.

----- Summary Statistics

Sequencing vector: plasmid; n/a; 100% of reads  
Chemistry: Dye-terminator Big Dye; 100% of reads  
Assembly program: Phrap; version 0.990319  
Consensus quality: 205824 bases at least Q40  
Consensus quality: 206436 bases at least Q30  
Consensus quality: 206755 bases at least Q20  
Insert size: 161000; agarose-fp  
Insert size: 206921; sum-of-contigs  
Quality coverage: 11.74x in Q20 bases; agarose-fp  
Quality coverage: 9.14x in Q20 bases; sum-of-contigs

\* NOTE: This is a 'working draft' sequence. It currently consists of 10 contigs. Gaps between the contigs are represented as runs of N. The order of the pieces is believed to be correct as given, however the sizes of the gaps between them are based on estimates that have been provided by the submitter.

\* This sequence will be replaced  
\* by the finished sequence as soon as it is available and the accession number will be preserved.

\* 1 22661: contig of 22661 bp in length  
\* 22662 22761: gap of unknown length  
\* 22762 45605: contig of 22844 bp in length  
\* 45606 45705: gap of unknown length  
\* 45706 64208: contig of 18503 bp in length  
\* 64209 70434: gap of unknown length  
\* 70435 70534: gap of unknown length  
\* 70535 123863: contig of 53329 bp in length  
\* 123864 123963: gap of unknown length  
\* 123964 133638: contig of 9675 bp in length  
\* 133639 133738: gap of unknown length  
\* 133739 151887: contig of 18149 bp in length  
\* 151888 151987: gap of unknown length  
\* 151988 161829: contig of 9842 bp in length  
\* 161830 161929: gap of unknown length  
\* 161930 181367: contig of 19438 bp in length  
\* 181368 181467: gap of unknown length  
\* 181468 207821: contig of 26354 bp in length.

#### FEATURES

##### Source

1..207821  
/organism="Bos taurus"  
/mol\_type="genomic DNA"  
/db\_xref="taxon:9913"  
/clone="RP42-405D15"  
/clone\_lib="RP42"

##### misc\_feature

1..22661  
/note="assembly\_fragment  
clone end:SP6  
vector side:left"

##### gap

22662..22761  
/estimated length=unknown

##### misc\_feature

22762..45605  
/note="assembly\_fragment"  
45606..45705

##### gap



```
/codon_start=1
/transl_table=11
/product="dtdp-4-keto-6-deoxyglucose-3,5-epimerase"
/protein_id="AADI9914.1"
/db_xref="GI:4406250"
/translation="MSDNFGKTLVVRKIDAI PGLLEFDIPVHGDRGWFKNPQKEK
MLPGFSPFPAAGKQNVSRKNGVRLGHAEPWPKYISVADDDGKVLGWSVDLREG
ETPGNVQTSIDKSGIFGPRVGANGFQVLSDTVSYSVLNDALELKPQYAFVNYA
DPALGHEWENLPEAEVSEADKHHLRLRDVXPLTKDEL"
3609.4655
/gene="cps19AN"
3609.4655
/gene="cps19AN"
/notes="Cps19AN"
/codon_start=1
/transl_table=11
/product="dtdp-glucose-4,6-dehydratase"
/protein_id="AADI9915.1"
/db_xref="GI:4406251"
/translation="MTEYKKIIVTGAGFGISNFVHYVNNFPDVHVTVLDKLTLYAGN
RANIEILGRVDELVDIADAALVDKLAADAI VHYAAESHNDNSLNDPSPFIHTN
FICTYLLAARXYDIFRHHVSTDEVVGLDLPLREDLPHGEGEGEKFARTKYNPSSP
YSSTKAADLIIVAWRSFGVKATISNCSNNYGPYQHIKEFIPROITNLSGKPKLY
GEGKVRDWHTHDHSQWTLITKGQIGETYLIGADGKNKEVELLELKGQPAD
AYHVTDRAGHDURYAIDASKLRDELGWKPEFTNFEAGLKETIKWYTDNQDWWKSEKE
AVEANYAKTQQVIK"
4659.4704
/complement(4717..5571)
/gene="cps19AO"
/complement(4717..5571)
/notes="Cps19AO"
/codon_start=1
/transl_table=11
/product="dtdp-L-rhamnose synthase"
/protein_id="AADI9916.1"
/db_xref="GI:4406252"
/translation="MLITGANGQLGTELRLNERNVDYVAVDVTMDITNSEMVEK
VFAEVPTLVHYCAATVADAEDGKELDFAINVGTENVAKSGKYGAILVYIGTD
FVDTGKPKVGOEVEDDI PDQTEYGRTRKMGELVENLTSQHYIIRTAWFGNYGRN
FYFTMQSLAETHKTLTVSDOYGRPTWTRTLAEPMYVTVENKEYGYHLSDAAEDT
TWDFASEILKESDVELLPVDSKFPKAKRPLNSTMSLAKAKATGFVPTWQDALKE
FYKQEVKK"
5817.>5832
/gene="alia"
5817.>5832
/gene="alia"
/codon_start=1
/transl_table=11
/product="Alia"
/protein_id="AADI9917.1"
/db_xref="GI:4406253"
/translation="MKSSK"
```

## ORIGIN

```
Query Match 24.0%; Score 31; DB 1; Length 5832;
Best Local Similarity 59.8%; Pred. No. 79;
Matches 52; Conservative 0; Mismatches 35; Indels 0; Gaps 0;

QY 32 ATTACTGACGACAAAGCCCGACCGAGATGGTGGGGTCTTTTGTGTGGTGTGTGA 91
Db 2722 ATTACAGACGTTAACCAAGGCTTACCTAAATCGTGGTGACCTTCTGTGAGCTGATGGG 2781
QY 92 CGGTGTGCCACCGTATTATCCGGA 118
Db 2782 CGTGGTTTCCGCTGGTGGATACGGA 2808
```

```
RESULT 14
CR931675
LOCUS CR931675 18617 bp DNA linear BCT 30-JUN-2005
DEFINITION Streptococcus pneumoniae strain Nr. 141/68 (serotype 19a).
ACCESSION CR931675
```

```
VERSION CR931675.1 GI:68643359
KEYWORDS Streptococcus pneumoniae
SOURCE Streptococcus pneumoniae
ORGANISM Bacteria; Firmicutes; Lactobacillales; Streptococcaceae; Streptococcus.
REFERENCE 1 (bases 1 to 18617)
AUTHORS Bentley,S.D., Aanensen,D., Mavroidi,A., Saunders,D., Rabinovitch,E., Collins,M., Donaghue,K., Harris,D., Kalltoft,M.S., Murphy,L., Quail,M.A., Samal,G., Skovsted,I.C., Barrell,B.G., Reeves,P., Parkhill,J. and Spratt,B.G.
TITLE Genetic analysis of the capsular biosynthetic locus from all 90 serotypes of Streptococcus pneumoniae
JOURNAL Unpublished
REFERENCE 2 (bases 1 to 18617)
AUTHORS Bentley,S.D.
TITLE Direct Submission
JOURNAL Submitted (09-DEC-2004) Submitted on behalf of the Pathogen Sequencing Unit, Sanger Institute, Wellcome Trust Genome Campus, Hinxton, Cambridge CB10 1SA, E-mail: edb@sanger.ac.uk
COMMENT NOTE: This sequence was generated from a PCR product representing the region from dexB to alia and is not necessarily responsible for the expressed capsule serotype. For a detailed description of how CUS products were predicted see the associated publication.
FEATURES
source
1..18617
/organism="Streptococcus pneumoniae"
/mol_type="genomic DNA"
/strain="Nr. 141/68"
/db_xref="taxon:1313"
<1..136
/gene="dexB"
/locus_tag="SPC19A_0001"
<1..136
/gene="dexB"
/locus_tag="SPC19A_0001"
/codon_start=2
/transl_table=11
/protein_id="CAI33621.1"
/db_xref="GI:68643360"
/translation="ANEQDLTVGKSVLIENTLAQEVFBKQILVPHWDAPFCVEMTD"
gene
complement(267..1697)
/gene="tnp"
/locus_tag="SPC19A_0002"
/pseudo
complement(267..1697)
/gene="tnp"
/locus_tag="SPC19A_0002"
/notes="member of homology group 0017 25 member of homology group 0017 25"
/pseudo
/codon_start=1
/transl_table=11
/product="putative IS1202 transposase"
1852..3297
/gene="wzg"
/locus_tag="SPC19A_0003"
1852..3297
/gene="wzg"
/locus_tag="SPC19A_0003"
/notes="member of homology group 0000 90"
/transl_table=11
/product="integral membrane regulatory protein Wzg"
/protein_id="CAI33621.1"
/db_xref="GI:68643361"
/translation="MSRRFKRSQKRSNVILTIYLLVCFLLFIKYNILAF
RYLNLVTVALLVALLIYKAEKFTIPLLVFSLVSSVLPVQOQVGLNLR
LNATSNYSVLSVAVLADSEIENVTLQMSVTPCTGDNIENQKLADIKSSQNTDLT
VNQSSYLAAYKSLIAGETKAIVLNSVFENIESEHPDYASKIKIYTKGPTKKVBP
KTSKQSFNIYVSGIDTGPISVSRSDNILMTVNRDTSKILLITTPRDADYVPIADG
GNNQKDKLTHAGIYGVDSIHTLENLYGVNINFTSFLKLDLGLGVDVYNDQ
```

```

DFTSLHGKHPFVGVNHLDSQALGFVRERYSLADGDHGRNQOKVIAAILQKLTS
EALKVSYRIIDSQDSIQTNMPLFTMINLNAQLESGTYKVNSQLKNGRDLPSY
AMPDSNLYMEINDSSLASVKTAIQDVLEGR"
/misc_feature
1852. .1968
/genes="wzg"
/locus_tag="SPC19A_0003"
/notes="Signal peptide predicted for SPC0963 by Signalp 2.0
HMM (Signal peptide probability 0.999) with cleavage site
probability 0.540 between residues 39 and 40"
join(1900. .1959,1987. .2055,2074. .2142)
/misc_feature
/genes="wzg"
/locus_tag="SPC19A_0003"
/notes="3 probable transmembrane helices predicted for
SPC0963 by TMHMM2.0 at aa 17-36, 46-68 and 75-97"
2059. .2403
/genes="wzg"
/locus_tag="SPC19A_0003"
/notes="HMPfam hit to PF02916, DNA polymerase processivity
factor, score 6.7e-65"
2581. .3024
/genes="wzg"
/locus_tag="SPC19A_0003"
/notes="HMPfam hit to PF03816, Cell envelope-related
transcriptional attenuator, score 2.8e-72"
3299. .4030
/genes="wzh"
/locus_tag="SPC19A_0004"
/notes="wzh"
3299. .4030
/genes="wzh"
/locus_tag="SPC19A_0004"
/EC_number="3.1.3.48"
/notes="member of homology group 0001 90"
/codon_start=1
/transl_table=11
/product="protein-tyrosine phosphatase Wzh"
/protein_id="CAI33624.1"
/db_xref="GI:68643362"
/translation="MIDIHSIHFVDVDDGPKSREESKALLAESVROGVRTIVSTSHR
KMFETPEEKIAENFLQVREIAKEVASDLVARGAEIYTTSDVDLKEKRIPTLNN
RYALIEFMTNTPYRDTHSALKIMLGITTVIAHIERYDALENNEKRVRLINNGCV
QVNSHYLKSGLFGERYKFMKCAQYFLERDLVHIITASDMHNVDSRPPHMAEAYDLVS
QKYGAEAKQSLFTDNPRIKIVMDQLI"
3611. .3925
/genes="wzh"
/locus_tag="SPC19A_0004"
/notes="HMPfam hit to PF02811, PHP, C-terminal, score
1e-11"
4036. .4731
/genes="wzd"
/locus_tag="SPC19A_0005"
4036. .4731
/genes="wzd"
/locus_tag="SPC19A_0005"
/notes="member of homology group 0002 90"
/codon_start=1
/transl_table=11
/product="capsular polysaccharide biosynthesis protein
wzd"
/protein_id="CAI33625.1"
/db_xref="GI:68643363"
/translation="MMKEONTIEIDVFQLLTKMKHLIILLVALVTGAGAPYSTFI
VKPEYSTRIVYVRNQGBKPLGTNQDLQNGTYLVKDYREIILSQDVLEKVTNLKL
DMFAKTUSKVQVTVPADTRIVSISVKDQGPASRIANSIREVAASKIIATVRSDV
TLLEEARPMTPESSPNRRTNLVGLGGAAVITVILLIELDTRVRRPDEVDELQVM
PLLGWVPDFDKMK"
4057. .4473
/genes="wzd"
/locus_tag="SPC19A_0005"
/notes="HMPfam hit to PF02706, Lipopolysaccharide
biosynthesis, score 1.6e-56"
join(4105. .4173,4573. .4641)
/genes="wzd"
/locus_tag="SPC19A_0005"

```

```

/notes="2 probable transmembrane helices predicted for
SPC0965 by TMHMM2.0 at aa 24-46 and 180-202"
4579. .4656
/genes="wzd"
/locus_tag="SPC19A_0005"
/notes="P800217 Sugar transport proteins signature 2."
4741. .5430
/genes="wze"
/locus_tag="SPC19A_0006"
4741. .5430
/genes="wze"
/locus_tag="SPC19A_0006"
/EC_number="2.7.1.112"
/notes="member of homology group 0003 90"
/codon_start=1
/transl_table=11
/product="tyrosine-protein kinase Wze"
/protein_id="CAI33626.1"
/db_xref="GI:68643364"
/translation="MPTLRISQAKELVKABEYYNALCTNLQSLGDDLKVFQITSYVK
AGEGKTTSTINAWARAGYKTLILDADNENSMVSGVKFSRERIITGLTFELSGITDL
SQGLCDINENLFIQAGSVSPNTALLQSKNFTMLEILRKYFDYLVVDTPAGVVII
DAAIMQKCDASILVTKAGETKRRLQKAKBQLEQTKSCLGVVLNKLDTVSVEKIGSY
GSYGNRYKQKK"
5445. .6806
/genes="wcha"
/locus_tag="SPC19A_0007"
5445. .6806
/genes="wcha"
/locus_tag="SPC19A_0007"
/notes="member of homology group 0005 66"
/codon_start=1
/transl_table=11
/product="undecaprenylphosphate glucosylphosphotransferase
Wcha (initial sugar transferase)"
/protein_id="CAI33627.1"
/db_xref="GI:68643365"
/translation="MSEKLAKPSVAIVQSPVLVILLTVLSAVRSTETIVSTTAVLYIL
HYFVYISDYGRNFKRGVLIJELVQTLKYLFPALASISNPFLEDRFSTSRGMIVF
LLHVFVLYVLPFIKWKRYAPNFKGSKILLLTATSRVEKVLDELISDDVVGSL
IDVTKDPDFQHDHDLKVAEGIVNFATHEVDEVFVNLPSEKINIGELVSPQETWG
IVTVNLNADFARNKQICEMAGLNVVTFSTTFYKTSHTVIAKRIIDVGLVGLILCG
LVSIVLPLIRKDGSAFQAQTRIGKNGRHFTYKFSKCMVDAEKKRELMQETMOG
GMFKVDDDPRIITKIGHFIRKTSLSDELQPVYNVLKDMSLVGTPTVDEYHTPEQK
RRLSFKPGITGLMQVSGRSEIKNFDEVKLDVAYIDGWTIMKDIELLTKVKVLMKD
GAK"
join(5469. .5528,5541. .5594,5628. .5696,5739. .5807,
6237. .6305)
/genes="wcha"
/locus_tag="SPC19A_0007"
/notes="5 probable transmembrane helices predicted for
Query Match 24.0%; Score 31; DB 1; Length 18617;
Best Local Similarity 59.8%; Pred. No. 78;
Matches 52; Conservative 0; Mismatches 35; Indels 0; Gaps 0;
Qy 32 ATTACTGACACAAAGACCCGACCGAGATGTCGGGGTCTTTTGTGTGTCGTGTA 91
||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 13947 ATTACAGACGTTAACRAGGCTTACCTAAATCGTGGTGACCTGATGGG 14006
||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Qy 92 CGTGTTCGCAACCGCTATTATTCGGGA 118
||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 14007 CGTGGTTTTCCTGGTGGATACGGGA 14033
||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
RESULT 15
AF094575 18754 bp DNA linear BCT 09-SEP-1999
LOCUS
DEFINITION Streptococcus pneumoniae serotype 19A dexB (dexB) gene, partial
sequence; capsular polysaccharide biosynthesis operon, complete
sequence; and oligopeptide permease Alia (alia) gene, partial cds.
AF094575
VERSION AF094575.1 GI:3907597

```

KEYWORDS

SOURCE Streptococcus pneumoniae  
ORGANISM Streptococcus pneumoniae  
Bacteria; Firmicutes; Lactobacillales; Streptococcaceae;  
Streptococcus.

REFERENCE

1 (bases 1 to 18754)  
Morona,J.K., Morona,R. and Paton,J.C.  
AUTHORS Analysis of the 5' portion of the type 19A capsule locus identifies  
TITLE two classes of cpsC, cpsD, and cpsE genes in Streptococcus  
pneumoniae

JOURNAL

J. Bacteriol. 181 (11), 3599-3605 (1999)

PUBLISHED

10348877

REFERENCE

2 (bases 1 to 18754)  
Morona,J.K., Morona,R. and Paton,J.C.  
AUTHORS Comparative genetics of capsular polysaccharide biosynthesis in  
TITLE Streptococcus pneumoniae types belonging to serogroup 19  
J. Bacteriol. 181 (17), 5355-5364 (1999)

JOURNAL

10464207

PUBLISHED

3 (bases 1 to 18754)

Morona,J.K., Morona,R. and Paton,J.C.  
AUTHORS Submitted (23-SEP-1998) Microbial Microbiology Unit, Women's and  
TITLE Children's Hospital, King William Rd, North Adelaide, SA 5006,  
JOURNAL Australia

FEATURES

source Location/Qualifiers

1..18754

/organism="Streptococcus pneumoniae"

/mol\_type="genomic DNA"

/serotype="19A"

/db\_xref="taxon:1313"

<1..9

/gene="dexB"

<1..9

/gene="dexB"

/codon\_start=1

/transl\_table=11

/product="DexB"

/protein\_id="AAC78662.1"

/db\_xref="GI:3907598"

/translation="VE"

complement(161..1647)

/note="insertion sequence; 3' truncated"

/rpt\_family="IS1202"

/rpt\_type=dispersed

1733..16682

/gene="capsular polysaccharide biosynthesis operon"

1733..3178

/gene="cps19a"

/gene="cps19a"

/codon\_start=1

/transl\_table=11

/product="putative regulatory protein Cps19a"

/protein\_id="AAC78663.1"

/db\_xref="GI:3907599"

/translation="MSRRFKSRQKVRNVNLTIVTLLVLCPLFLIFTYNILAF  
RYNLVLPALVLLIILYKAEKFTIPLVFSVSVLFAVQQFGLTNR  
LNATSVYSIVAVLADSEIENVTLMSVAPTGTDNENIOLKADIKSSONTOLT  
NOSSVLAAYKSLIAGETKAVLVNSVFENIIESEHPDYSKIKIYTKGTFKVEAP  
KTSKDSFNIVSGIDTYGPISSVSRSYNIMTVNRDTKILLITTPRDAYPIADG  
GNQKDLTHAGIYGVDSSTHTUENLYGVNDINTYRNLFTSFUKLIDLLGGVDVINDQ  
DFTSLGKHPFVGNVHLDSQALGFVRERYSLADGDHGRNQKQVIAAILQKLTSS  
EALKNTSRIIDSLQDSIQTNMPLTMINLVNALESGGTYKVNSQLKGNRTDLPST  
AMPDSNLYMEINDSSLASVKTAIQDVLGR"

3180..3911

/gene="cps19aB"

/gene="cps19aB"

/codon\_start=1

/transl\_table=11

/product="Cps19aB"

/protein\_id="AAC78664.1"

/db\_xref="GI:3907600"

gene

CDS

/translation="MIDIHSHIVFDVDDGPKGREESKALLAESYRQGVRTIVTSTSHRR  
KGMFETPEEKIAENFLQVRSIAKEVASDLVIAYGAEIYYTSDVLDKLEKNRIFTLNNS  
RYALIEFNTPTYRDIHSALSILMIGTIPVIAHIERYDALENNEKRVRELINMGCVT  
QVNSHVLSKSLFGERYKPMKRAQYFLERDLVHIITASDMHNVDSRPHPMAEYDLVS  
QVYGEAKAQLFIDNPRKIVMDQLI"

gene

CDS

3917..4612

/genes="cps19aC"

3917..4612

/genes="cps19aC"

/function="chain length regulation and export"

/codon\_start=1

/transl\_table=11

/product="Cps19aC"

/protein\_id="AAC78665.1"

/db\_xref="GI:3907601"

/translation="MMKEQNTIEIDVFQLLKTILWKHKLIIILVALVTGAGAFAYSTFI  
VKPEYTSITRIYVNRNQDKPLTNQDLQAGTYLVKDYREIILSDVLSKVAITNLKL  
DMPAKTLTSKVQVTPADTRIVSIVKDKQPEASRIANSLEVAEAKIINAVTRSDV  
TTLEEARPANTFSSPNVRRNTLVGLGGAATVITVILLIELLDTRVCRPEDVEDVLQM  
PLLVVVPDFDKMK"

gene

CDS

4622..5311

/genes="cps19aD"

4622..5311

/genes="cps19aD"

/function="chain length regulation and export"

/codon\_start=1

/transl\_table=11

/product="Cps19aD"

/protein\_id="AAC78666.1"

/db\_xref="GI:3907602"

/translation="MPTLSIQAKLELVKAEYYNALCTNLQSGDDLKVFISITSVK  
AGEGKTTSTNIAPARAGYKTLIDADMNSVMSGVFKSRERITGLTFELSGTTDL  
SQGLCDTNIENLFVQAGSVSPNTALQSKNPTTLMLETRKYFDYIVVDVAPGVVI  
DAAIIMQKDSASILVTKAGETKRDLQKAQLEQTKGKSLGVVLNKLDTSEVKYGSY  
GSVGYNRKOKX"

gene

CDS

5326..6687

/genes="cps19aE"

5326..6687

/genes="cps19aE"

/codon\_start=1

/transl\_table=11

/product="glucosyl-1-phosphate transferase Cps19aE"

/protein\_id="AAC78667.1"

/db\_xref="GI:3907603"

/translation="MSEKLAKPSVAIVQSFVLVLTLLYSAVRETIIVTAVLXIL  
HYFVYISDYGRNFKRGYLIBLVOTLKILFALAISNPFLEDFSIISRRGMIFY  
LILHVLVTVLNPFIKWYKRAYPNPKGSKKILLLTATSRVKVLDRLLIESDDVCEL  
VAVSLDKPFDQHDLLKVVAEGEIVNPATHEVDEVFINLPSEKYNIGELVSQFETMG  
IDVTNLNAPDWARNKQICEMAGLNVVTFSTTFYKTSHVIAKRIIDVLSGLVLICG  
LVSIVLPLIRKDGSAFPAQTRIGKNGRHFTFYKPSMCDABEKKRELMQNTWQG  
GMFKVDDDPRIITKIGHFIRKTSIDELPQFYNVLKGDMSLVGTRPPTVDEHYHTPBQK  
RLSPKPGITGLWQVSGRSEIKNFDEVKLDVAYIDGMTIMKDIETLLKTVKVLMDK  
GAK"

gene

CDS

6693..7436

/genes="cps19aF"

6693..7436

/genes="cps19aF"

/codon\_start=1

/transl\_table=11

/product="putative N-acetyl-mannosamine transferase  
Cps19aF"

/protein\_id="AAC78668.1"

/db\_xref="GI:3907604"

/translation="MNERIQILGITIDPLTMKETVDAVEQYVLKXHLPLHLMGVNADKI  
NOCYDEKIKKIVNESGIIINADGSAVLASKPLGTVPVPRVACIDIMKHLLELSNKG  
YSVYFGAKEDVLQDMLKIFPEERYPALNVGYRNGYFSEDEKHIQBDIKKKPDPVF  
VGITSPKSYIIIQSFMDNGINAVFMGGSGFDVLSGHIKRAPLMMQNAHLEWLFVRAN  
EPKRLFKRYFVGNVSFIRKVLRAKRGYKI"

gene

CDS

7475..8257

/genes="cps19aG"

7475..8257

/genes="cps19aG"

/function="putative protein involved in formation of



```
phospho-diester linkage"
/codon_start=1
/transl_table=11
/product="cps19aJ"
/protein_id="AAC78669.1"
/db_xref="GI:3907605"
/translation="MDAIEKFEKICKENNIDFFLRGSSVLGAVKYGDFIPWDDMDIA
VPREGYDKLPGIKFDRILAGKYQVLAYQCDTHCYEPRLFLLERKRLGLPRNTNL
GLHLIDIIPLDGAPNHSFLKLYFGKVIYWRFLASLGTITVGDHVMHSAKQKLIIGF
FKLGGFAKLPQNYVYKRDNLKDYKDKWKKQSLTINASLFAKEVMPVEIWEGVEK
PFEDAFKVPTEYDRYLKRLYGENYLHEEPSDDDKKSHLGGQ"
gene      8259..9137
CDS       8259..9137
          /gene="cps19aH"
          /gene="cps19aH"
          /codon_start=1
          /transl_table=11
          /product="putative rhamnosyl transferase Cps19aH"
          /protein_id="AAC78670.1"
          /db_xref="GI:3907606"
          /translation="MFCYIIHLYKVLBEETISCVKSIKEGNSNTKQIVIIDFNSNGTG
          EKLQELYESDEIDVLINHENAGFANNVAYQFAKEKYKDPFMWIMNNDIELETEEF
          EKIYDIIYRKEKHLGPDIFSTYQLHQNPRLTHYTYEEVKALNEKFKKQSQVSLA
          LKTKWLKSSKVLRTAIYQNRKKKSVDYRKQVENPILHGSFIVYSRDLIEKEEYAFN
          PNTFFVYETILDYEARLKGCKRIYTPKIKVLHHQNVAITNQVYTNLVEKTLFSNKNCF
          ESTSYFLKLNKENG"
          9138..10472
          /gene="cps19aI"
          9138..10472
          /gene="cps19aI"
          /codon_start=1
          /transl_table=11
          /product="polysaccharide polymerase Cps19aI"
          /protein_id="AAC78671.1"
          /db_xref="GI:3907607"
          /translation="MTYFLLLCLTLFLTFPFYFAFNQDLIAPPVVMFLISSVPA
          LINQWNIYSGLAYLIIISGLIVFSMPILLNLSPLNTKIKVTDRLIDIQFWKIAL
          TIIVDLIIILYRREIHNLAISHGYTGSNFQWFFRNATSYEGELTVRTSIRVLRIID
          VSAIYFGTYTINFFIYSHKRSKDLLVLPFLIFISKLLSGGRLLDIIKLLIAYVNA
          YIQQRKVGWDKVIISHKYMRLGTFVGLIAGIPTFYISLFSRSTRTVFESISTYLGQ
          SIQHFNQYIQNPICVAEYFGDESFAIMNIIKGLNLFVYVNSTVHLEFRQLGITMGVY
          TFFRRPHWDGLGVMYIFSPVGVFFAIFYLKLKRSRAGFKLDIHTIITYSYFYWIPL
          SSIEQXSFTTISLFTLVFIVLVYLMAFYWNLDHFHRGKLVKLSDTSIKSEN"
          10497..11924
          /gene="cps19aJ"
          10497..11924
          /gene="cps19aJ"
          /codon_start=1
```

```
Query Match      24.0%; Score 31; DB 1; Length 18754;
Best Local Similarity 59.8%; Pred.No. 78;
Matches 52; Conservative 0; Mismatches 35; Indels 0; Gaps 0;

Qy      32  ATTACTGACGACGAAGACCCGACGAGATGGTCGGGTCCTTTTGTGTGGTCTGTGA 91
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db      13828 ATTACAGCGTTACACAGGCTTACCTAAATCGTGTGACCTTCTGTTGAGCTGATGGG 13887

Qy      92  CGTGTTCCTCAACCGTATTATTCGGA 118
      ||||| ||||| ||||| |||||
Db      13888 CGTGGTTTTCGCTGGTTGGATACGGA 13914
```

Search completed: March 7, 2006, 12:15:11  
Job time : 1202.77 secs



GenCore version 5.1.7  
Copyright (c) 1993 - 2006 Bioceleration Ltd.

OM nucleic - nucleic search, using sw model

Run on: March 7, 2006, 10:55:04 ; Search time 304.617 Seconds  
(without alignments)  
2822.383 Million cell updates/sec

Title: US-10-782-899-1\_COPY\_472\_600

Perfect score: 129  
Sequence: 1 tgacctctgctgtagcga.....tattccgactagttcagcg 129

Scoring table: IDENTITY NUC  
Gapop 10.0 , Gapext 1.0

Searched: 4996997 segs, 3332346308 residues

Total number of hits satisfying chosen parameters: 9993994

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%  
Listing first 45 summaries

Database : N Geneseq\_21.\*  
1: Geneseqn1980s.\*  
2: Geneseqn1980s.\*  
3: Geneseqn2000s.\*  
4: Geneseqn2001as.\*  
5: Geneseqn2001bs.\*  
6: Geneseqn2002as.\*  
7: Geneseqn2002bs.\*  
8: Geneseqn2003as.\*  
9: Geneseqn2003bs.\*  
10: Geneseqn2003cs.\*  
11: Geneseqn2003ds.\*  
12: Geneseqn2004as.\*  
13: Geneseqn2004bs.\*  
14: Geneseqn2005s.\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
1	129	100.0	600	ABK52324	Abk52324 DNA encod
c	129	100.0	156638	6 ABQ81850	Abq81850 Bifidobac
3	30.6	23.7	26309	4 AAS59535	Aas59535 Propionib
4	30.6	23.7	26309	8 ACF64464	Acf64464 Propionib
5	29.6	22.9	2453	10 ADE63675	Ade63675 Human gen
6	29.4	22.8	870	13 ADK44809	Adk44809 Streptoco
7	29.4	22.8	885	13 ADR91671	Adr91671 Novel S.
8	29.4	22.8	885	14 AEA55541	Aea55541 Streptoco
c	29.4	22.8	1377	12 ADQ89190	Adq89190 Non-natur
10	29.4	22.8	3520	5 ADL45225	Adl45225 Human ova
c	29.2	22.6	634	6 ABL01350	AbL01350 Murine ap
12	29.2	22.6	1659	14 ADM69564	Adm69564 DNA encod
13	29.2	22.6	1760	11 AEA79104	Aea79104 Human apo
14	29.2	22.6	1800	13 ADR89493	Adr89493 Apoptosis
15	29.2	22.6	1836	6 ABL01352	AbL01352 Murine ap
16	29	22.5	110000	14 AEA61160_0	Aea61160 Human NED
c	28.6	22.2	1146	11 ABD03998	Abd03998 Pseudomon
18	28.6	22.2	2184	11 ABD03637	Abd03637 Pseudomon
19	28.6	22.2	2868	11 ABD03707	Abd03707 Pseudomon

20	28.4	22.0	936	2 AAT79329	Aat79329 DNA encod
21	28.4	22.0	936	8 ABX11507	Abx11507 DNA encod
22	28.4	22.0	936	10 AAD58914	Aad58914 Archaeogl
23	28.4	22.0	2451	6 AB212259	Ab212259 Arabidops
c	28.2	21.9	1152	5 AAS73582	Aas73582 DNA encod
24	28.2	21.9	1152	12 ACH79902	Ach79902 Human gen
25	28.2	21.9	1152	5 AAS73582	Aas73582 DNA encod
26	28.2	21.9	13540	8 ABZ74138	Abz74138 Secreted
27	28.2	21.9	13540	8 ADA98673	Ada98673 Human sec
28	28.2	21.9	13540	10 ABT16951	Abt16951 Human sec
29	28.2	21.9	13540	10 ABZ67695	Abz67695 Human sec
c	28.2	21.9	18038	5 ADM20219	Adm20219 Alternati
30	28.2	21.9	18038	8 ABZ74137	Abz74137 Secreted
31	28.2	21.9	18038	8 ADA98672	Ada98672 Human sec
32	28.2	21.9	18038	10 ABT16950	Abt16950 Human sec
33	28.2	21.9	18038	10 ABZ67694	Abz67694 Human sec
34	28.2	21.9	18038	10 ACN44524	Acn44524 Mouse gen
c	28	21.7	165221	11 ADZ13004	Adz13004 Murine ca
36	28	21.7	165799	14 AAS46220	Aas46220 Human DNA
c	27.8	21.6	2037	4 ACA89670	ACA89670 cDNA enco
38	27.8	21.6	2037	8 ACA73680	ACA73680 Human sec
c	27.8	21.6	2037	8 ACA05995	ACA05995 Human sec
c	27.8	21.6	2037	8 ACA66829	ACA66829 cDNA enco
c	27.8	21.6	2037	8 ACF20404	ACF20404 Human sec
c	27.8	21.6	2037	8 ACF19790	ACF19790 Human sec
c	27.8	21.6	2037	8 ACD22078	ACD22078 Human sec
c	27.8	21.6	2037	8 ACF13243	ACF13243 Human sec

ALIGNMENTS

RESULT 1  
ABK52324  
ID ABK52324 standard; DNA; 600 BP.  
XX  
AC ABK52324;  
XX  
DT 13-AUG-2002 (first entry)  
XX  
DE DNA encoding cancer gene therapy associated protein.  
XX  
KW Solid cancer; anaerobic bacteria; antitumour active protein; cancer;  
KW gene therapy; gene; ds.  
XX  
OS Bifidobacterium longum.  
XX  
FH Key Location/Qualifiers  
FT CDS 193..474  
FT /\*tag= a  
FT /product= "Cancer gene therapy protein"  
XX  
PN JP2002097144-A.  
XX  
PD 02-APR-2002.  
XX  
PF 21-SRP-2001; 2001JP-00290187.  
XX  
PR 21-SRP-2000; 2000JP-00287688.  
XX  
PA (AMAN/) AMANO A.  
PA (FUJI/) FUJIMORI M.  
DR WPI; 2002-448201/48.  
DR P-PSDB; AAU96807.  
XX  
PT Solid cancer therapy with anaerobic bacteria of Bifidobacterium sp. by  
PT tumor tissue specific delivery of a DNA encoding for an antitumor active  
PT protein or its precursor.  
XX  
PS Claim 10; Page 16; 21pp; Japanese.  
XX  
CC The invention describes a method of treating a solid cancer with  
CC anaerobic bacteria by site specific delivery of DNA encoding an



CC associated DNA sequences are used in the treatment, prevention and  
 CC diagnosis of medical conditions caused by P. acnes. The disorders include  
 CC SAPHO syndrome (synovitis, acne, pustulosis, hypertonosis and  
 CC osteomyelitis), uveitis and endophthalmitis. P. acnes is also involved in  
 CC infections of bone, joints and the central nervous system, however it is  
 CC particularly involved in the inflammatory lesions associated with acne  
 CC vulgaris. A method for detecting the presence or absence of P. acnes in a  
 CC patient comprises contacting a sample with a binding agent that binds to  
 CC the proteins of the invention and determining the amount of bound protein  
 CC in the sample. The polypeptides may be used as antigens in the production  
 CC of antibodies specific for P. acnes proteins. These antibodies can be  
 CC used to downregulate expression and activity of P. acnes polypeptides and  
 CC therefore treat P. acnes infections. The antibodies may also be used as  
 CC diagnostic agents for determining P. acnes presence, for example, by  
 CC enzyme linked immunosorbent assay (ELISA). This sequence encodes the  
 CC polypeptides shown in AAU46704-AAU46985 and AAU67509. Note: The sequence  
 CC data for this patent did not form part of the printed specification, but  
 CC was obtained in electronic format directly from WIPO at  
 CC ftp.wipo.int/pub/published\_pct\_sequences  
 XX  
 SQ Sequence 26309 BP; 5546 A; 8173 C; 7885 G; 4699 T; 0 U; 6 Other;  
 Query Match 23.7%; Score 30.6; DB 4; Length 26309;  
 Best Local Similarity 58.1%; Pred. No. 11;  
 Matches 54; Conservative 0; Mismatches 39; Indels 0; Gaps 0;  
 QY 37 TGACGACAAAGACCCCGACGAGATGTCGGGCTCTTTTGTGTGCTGTGACGTGT 96  
 Db 9538 TGACGACGACCATCCCGACTCCCATATTGAGGGTCGCTTCGATGTCAGGCTGTGACAGCT 9597  
 QY 97 TGTCCAAACCGTATTATTCGGACTAGTTTCAGCG 129  
 Db 9598 TGCCCCACCTGCTGGACGAGGTTGAAGATCGCG 9630  
 RESULT 4  
 ACF64464  
 ID ACF64464 standard; DNA; 26309 BP.  
 XX  
 AC ACF64464;  
 XX  
 DT 17-OCT-2003 (first entry)  
 XX  
 DE Propionibacterium acnes DNA contig sequence #30.  
 XX  
 KW Acne vulgaris; antiseborrheic; dermatological; antibacterial;  
 KW immunostimulant; immune response; vaccine; ds.  
 XX  
 OS Propionibacterium acnes.  
 XX  
 PN WO2003033515-A1.  
 XX  
 PD 24-APR-2003.  
 XX  
 PF 11-OCT-2002; 2002WO-US032727.  
 XX  
 PR 15-OCT-2001; 2001US-00978825.  
 XX  
 PA (CORI-) CORIXA CORP.  
 XX  
 PI Mitcham JL, Skeiky YAW, Persing DH, Bhatia A, Maisonneuve JL;  
 PI Zhang Y, Wang S, Jen S, Lodes MJ, Benson DR, Jones R, Carter D;  
 PI Barth B, Valliave-Douglas J;  
 XX  
 DR WPI; 2003-381789/36.  
 XX  
 PT New Propionibacterium acnes polypeptides and polynucleotides encoding the  
 PT polypeptide, useful for diagnosing, preventing or treating acne vulgaris,  
 PT or for stimulating an immune response specific for a P. acnes protein.  
 XX  
 PS Claim 1; SEQ ID NO 30; 1481pp; English.  
 XX  
 CC The invention relates to an isolated polynucleotide (ACF64435-ACF64733)

CC encoding a Propionibacterium acnes protein. The invention also relates to  
 CC polypeptides encoded by the polynucleotides (ABM35624-ABM4536) and to  
 CC immunogenic fragments of P. acnes polypeptides. The invention  
 CC additionally encompasses expression vectors and host cells comprising a  
 CC polynucleotide of the invention; antibodies against polypeptides of the  
 CC invention; fusion proteins comprising a polypeptide of the invention; a  
 CC method for stimulating an immune response specific for a P. acnes  
 CC polypeptide and an isolated T cell population comprising T cells prepared  
 CC via this method; a vaccine composition (comprising P. acnes polypeptides,  
 CC polynucleotides, antibodies, fusion proteins, T cell populations, or  
 CC antigen-presenting cells that express the polypeptide); a method and kit  
 CC for detecting or determining the presence or absence of P. acnes in a  
 CC patient; and a method for inhibiting the development of P. acnes in a  
 CC patient. The P. acnes polypeptides, polynucleotides, antibodies, fusion  
 CC proteins, T cell populations or antigen-presenting cells that express the  
 CC polypeptides are useful for diagnosing, preventing or treating acne  
 CC vulgaris, or for stimulating an immune response specific for a P. acnes  
 CC protein. The polynucleotides can also be used as probes or primers for  
 CC nucleic acid hybridisation. The vaccine composition is useful for the  
 CC stimulation of an immune response against P. acnes, or for treating acne,  
 CC and the kit is useful for performing a diagnostic assay. The present  
 CC sequence represents a P. acnes DNA contig which is specifically claimed  
 CC in the invention. Note: The sequence data for this patent did not form  
 CC part of the printed specification, but was obtained in electronic format  
 CC directly from WIPO at ftp.wipo.int/pub/published\_pct\_sequences  
 XX  
 SQ Sequence 26309 BP; 5546 A; 8173 C; 7885 G; 4699 T; 0 U; 6 Other;  
 Query Match 23.7%; Score 30.6; DB 8; Length 26309;  
 Best Local Similarity 58.1%; Pred. No. 11;  
 Matches 54; Conservative 0; Mismatches 39; Indels 0; Gaps 0;  
 QY 37 TGACGACAAAGACCCCGACGAGATGTCGGGCTCTTTTGTGTGCTGTGACGTGT 96  
 Db 9538 TGACGACGACCATCCCGACTCCCATATTGAGGGTCGCTTCGATGTCAGGCTGTGACAGCT 9597  
 QY 97 TGTCCAAACCGTATTATTCGGACTAGTTTCAGCG 129  
 Db 9598 TGCCCCACCTGCTGGACGAGGTTGAAGATCGCG 9630  
 RESULT 5  
 ADE63675  
 ID ADE63675 standard; DNA; 2453 BP.  
 XX  
 AC ADE63675;  
 XX  
 DT 29-JAN-2004 (first entry)  
 XX  
 DE Human gene NM\_004046, SEQ ID NO 9619.  
 XX  
 KW Human; ds; gene; pain; neuronal tissue; gene therapy;  
 KW spinal segmental nerve injury; chronic constriction injury; CCI;  
 KW spared nerve injury; SNI; Chung.  
 XX  
 OS Homo sapiens.  
 XX  
 PN WO2003016475-A2.  
 XX  
 PD 27-FEB-2003.  
 XX  
 PF 14-AUG-2002; 2002WO-US025765.  
 XX  
 PR 14-AUG-2001; 2001US-0312147P.  
 PR 01-NOV-2001; 2001US-0346382P.  
 PR 26-NOV-2001; 2001US-0333347P.  
 XX  
 PA (GEHO ) GEN HOSPITAL CORP.  
 PA (FARB ) BAYER AG.  
 XX  
 PI Woolf C, D'urso D, Befort K, Costigan M;  
 XX  
 DR WPI; 2003-368312/36.

DR GENBANK; NM\_004046.  
XX  
PT New composition comprising two or more isolated polypeptides, useful for  
PT preparing a medicament for treating pain in an animal.  
XX  
PS  
PS Claim 1; Page; 1017pp; English.  
XX  
XX The invention discloses a composition comprising two or more isolated rat  
CC or human polynucleotides or a polynucleotide which represents a fragment,  
CC derivative or allelic variation of the nucleic acid sequence. Also  
CC claimed are a vector comprising the novel polynucleotide, a host cell  
CC comprising the vector, a method for identifying a nucleotide sequence  
CC which is differentially regulated in an animal subjected to pain and a  
CC kit to perform the method, an array, a method for identifying an agent  
CC that increases or decreases the expression of the polynucleotide sequence  
CC that is differentially expressed in neuronal tissue of a first animal  
CC subjected to pain, a method for identifying a compound which regulates  
CC the expression of a polynucleotide sequence which is differentially  
CC expressed in an animal subjected to pain, a method for identifying a  
CC compound that regulates the activity of one or more of the  
CC polynucleotides, a method for producing a pharmaceutical composition, a  
CC method for identifying a compound or small molecule that regulates the  
CC activity in an animal of one or more of the polypeptides given in the  
CC specification, a method for identifying a compound useful in treating  
CC pain and a pharmaceutical composition comprising the one or more  
CC polypeptides or their antibodies. The polynucleotide or the compound that  
CC modulates its activity is useful for preparing a medicament for treating  
CC pain (e.g. spinal segmental nerve injury (Chung), chronic constriction  
CC injury (CCI) and spared nerve injury (SNI)) in an animal (e.g. gene  
CC therapy). The sequence presented is a human DNA (shown in Table 2 of the  
CC specification) which encodes one of the polypeptides of the invention  
CC which is differentially expressed during pain. Note: The sequence data  
CC for this patent did not form part of the printed specification, but was  
CC obtained in electronic form directly from WIPO at  
CC ftp.wipo.int/pub/published\_pct\_sequences.  
XX  
SQ Sequence 2453 BP; 622 A; 536 C; 635 G; 660 T; 0 U; 0 Other;  
  
Query Match 22.9%; Score 29.6; DB 10; Length 2453;  
Best Local Similarity 57.6%; Pred. No. 12;  
Matches 53; Conservative 0; Mismatches 39; Indels 0; Gaps 0;  
  
QY 4 CCTTCTGCTGCTAGCGATTCTCGAGCATTACTGACGACAAAGACCCGACCGAGATGG 63  
DB 1864 CCATCACTCGGACAGATTCTTGGACACAGATTGCTACAAAGATCCGCCCTCAATTA 1923  
  
QY 64 TCGGGGTCCTTTTGTGTGTGCTGTGACGTG 95  
DB 1924 ACGTGGTCTGCTCGTATCTCGTTCGGATCCGTG 1955  
  
RESULT 6  
ADK44809  
ID ADK44809 standard; DNA; 870 BP.  
XX  
AC ADK44809;  
XX  
XX 24-FEB-2005 (first entry)  
XX  
XX Streptococcus pneumoniae gene, Seq ID No 1324.  
XX  
XX ds; gene; Antibacterial; Gene therapy; Vaccine; Streptococcus pneumoniae.  
XX  
OS Streptococcus pneumoniae.  
XX  
XX US6699703-B1.  
XX  
XX 02-MAR-2004.  
XX  
XX 26-MAY-2000; 2000US-00583110.  
XX  
XX 02-JUL-1997; 97US-0051553P.  
XX  
XX 12-MAY-1998; 98US-0085131P.  
XX

PR 30-JUN-1998; 98US-00107433.  
XX  
PA (GENO-) GENOME THERAPEUTICS CORP.  
XX  
PI Doucette-Stamm L, Bush D, Zeng Q, Opperman T, Housewheat CE;  
XX  
XX WPI; 2004-212399/20.  
DR P-PSDB; ADK47470.  
XX  
XX New nucleic acid molecules and polypeptides useful for diagnosing,  
XX PT preventing and treating pathological conditions resulting from bacterial  
PT infection, e.g. Streptococcus pneumoniae infection, and in drug  
PT screening.  
XX  
XX Disclosure; SEQ ID NO 1324; 301pp; English.  
PS  
XX The invention relates to isolated Streptococcus pneumoniae nucleic acids  
CC and polypeptides. The nucleic acids and proteins are useful for  
CC diagnosing, preventing and treating pathological conditions resulting  
CC from bacterial infection, such as S. pneumoniae infection. These may also  
CC be used for drug screening procedures. The present sequence represents a  
CC Streptococcus pneumoniae nucleic acid of the invention. Note: The  
CC sequence data for this patent did not appear in the printed specification  
CC but was obtained in electronic format directly from USPTO at  
CC seqdata.uspto.gov/sequence.html.  
XX  
SQ Sequence 870 BP; 237 A; 166 C; 215 G; 252 T; 0 U; 0 Other;  
  
Query Match 22.8%; Score 29.4; DB 13; Length 870;  
Best Local Similarity 58.6%; Pred. No. 10;  
Matches 51; Conservative 0; Mismatches 36; Indels 0; Gaps 0;  
  
QY 32 ATTACTGACGACAAAGACCCGACCGAGATGGTCGGGTCCTTTTGTGTGCTGTGTA 91  
DB 589 ATTACAGATATCAACAAAGCGCTTACCTAAATCGTGGTGACCTTCTGTGAGCTGATGGG 648  
  
QY 92 CGTGTGTGTCACACCGTATTATTCGGA 118  
DB 649 CGTGTGTGTCGTTGGTACGGA 675  
  
RESULT 7  
ADR91671  
ID ADR91671 standard; DNA; 885 BP.  
XX  
AC ADR91671;  
XX  
XX 16-DEC-2004 (first entry)  
XX  
XX Novel S. pneumoniae DNA sequence, SEQ ID 306.  
XX  
XX Meningitis; bacteraemia; pneumonia; otitis media; ds;  
XX bacterial infection.  
XX  
OS Streptococcus pneumoniae.  
XX  
XX US6800744-B1.  
XX  
XX 05-OCT-2004.  
XX  
XX 30-JUN-1998; 98US-00107433.  
XX  
XX 02-JUL-1997; 97US-0051553P.  
XX  
XX 12-MAY-1998; 98US-0085131P.  
XX  
XX (GENO-) GENOME THERAPEUTICS CORP.  
XX  
XX Doucette-Stamm LA, Bush D;  
XX  
XX WPI; 2004-697205/68.  
DR P-PSDB; ADR94274.  
XX  
XX New isolated nucleic acid encoding a Streptococcus pneumoniae  
PT









```
XX DE Human apoptosis-associated cDNA SEQ ID 209.
XX
XX apoptosis; cell death; cytostatic; neuroprotective; immunosuppressive;
XX anti-rheumatic; antiarthritic; dermatological; anti-inflammatory;
XX hepatotropic; virucide; nootropic; anticonvulsant; antiparkinsonian;
XX vasotropic; cerebroprotective; antialcoholic; gene therapy; tumour;
XX autoimmune disease; degenerative disease; viral infection; leukaemia;
XX carcinoma; sarcoma; multiple sclerosis; rheumatoid arthritis; diabetes;
XX lupus; hepatitis; influenza viruses; Alzheimer's disease;
XX Huntington's disease; Parkinson's disease; reperfusion injury; stroke;
XX alcoholic liver disease; human; gene; ss.
XX
XX Homo sapiens.
XX
XX WO2003058021-A2.
XX
XX 17-JUL-2003.
XX
XX 13-JAN-2003; 2003WO-EP000270.
XX
XX 11-JAN-2002; 2002DE-01000856.
XX
XX (XANT-) XANTOS BIOMEDICINE AG.
XX
XX Koenig-Hoffman K, Kazinski M, Schaefer R, Kesper B;
XX
XX WPI; 2003-542134/51.
XX
XX New nucleic acids involved in apoptosis, useful for diagnosis and
XX treatment of e.g. tumors and degenerative disease, also related proteins,
XX antibodies and modulators.
XX
XX Claim 1a; SEQ ID NO 209; 517pp; German.
XX
XX This invention describes novel nucleic acid molecules that are associated
XX with apoptosis and encode a polypeptide and are derived from a normalised
XX gene library (embryonic or liver) or clone collections, and the extent of
XX apoptosis measured by cell death detection assay or the CPRG assay
XX (measuring loss of membrane integrity). The products of the invention
XX have cytostatic, neuroprotective, immunosuppressive, anti-rheumatic,
XX antiarthritic, dermatological, anti-inflammatory, hepatotropic, virucide,
XX neurotropic, anticonvulsant, antiparkinsonian, vasotropic,
XX cerebroprotective and antialcoholic activity and can be used for gene
XX therapy. The polynucleotides also related vectors, hosts (or their
XX extracts), encoded polypeptide (or their receptors) and/or agents that
XX inhibit their activity (including antisense sequences) are used for
XX treatment or prevention of tumours, autoimmune or degenerative diseases
XX and viral infections, specifically leukaemia, carcinoma, sarcoma,
XX multiple sclerosis, rheumatoid arthritis, diabetes, lupus, or infection
XX with hepatitis or influenza viruses, Alzheimer's, Huntington's or
XX Parkinson's disease, reperfusion injury, stroke and alcoholic liver
XX disease. Detection of the polynucleotides and derived polypeptides can
XX also be used for diagnosis of these diseases. This sequence encodes an
XX apoptosis-associated protein described in the disclosure of the
XX invention.
XX
XX Sequence 1760 BP; 326 A; 513 C; 535 G; 386 T; 0 U; 0 Other;
XX
XX Query Match 22.6%; Score 29.2; DB 11; Length 1760;
XX Best Local Similarity 59.8%; Pred. No. 15;
XX Matches 49; Conservative 0; Mismatches 33; Indels 0; Gaps 0;
XX
XX Qy 48 ACCCGACCGAGATGGTGGGGTCTTTTGTGTGGTGTGTGACGTGTGTCCACCGT 107
XX | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
XX Db 762 ATCCCTCACCTCCTGGGGGATGGTGTGTGTGGGCTGTAACTGCTGCCCACTTC 821
XX | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
XX Qy 108 ATTATTCGGGACTAGTTCAGCG 129
XX | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
XX Db 822 ATCAATGCCTACTTGGTGGACG 843
XX | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
XX
XX RESULT 14
XX ID ABL01352 standard; DNA; 1836 BP.
XX
XX ABL01352;
XX
XX 15-MAR-2002 (first entry)
```

```
ADR89493
ID ADR89493 standard; DNA; 1800 BP.
XX
XX ADR89493;
AC
DT 02-DEC-2004 (first entry)
XX
DE Apoptosis-inducing protein coding sequence, SEQ ID 17.
XX
XX Cytostatic; Immunosuppressive; Neuroprotective; Vasotropic; Virucide;
XX apoptosis; neurodegenerative disease; ischaemic disease; cancer;
XX autoimmune disease; viral disease; murine; gene; ds.
XX
XX Mus musculus.
XX
XX Key Location/Qualifiers
XX CDS 1..1170
XX FT /*tag= a
XX FT /product= "Apoptosis-inducing protein"
XX
XX WO2004078112-A2.
XX
XX 16-SEP-2004.
XX
XX 05-MAR-2004; 2004WO-JP002899.
XX
XX 07-MAR-2003; 2003JP-00061179.
XX
XX 10-MAR-2003; 2003US-0452943P.
XX
XX (ASAH-) ASAHI KASEI PHARMA CORP.
XX
XX Muramatsu S, Takeda M, Matsuoka A;
XX
XX WPI; 2004-662343/64.
XX
XX P-PSDB; ADR89494.
XX
XX New protein capable of inducing apoptosis, useful in screening for
XX compounds that inhibit or induce apoptosis which may be used to treat
XX neurodegenerative, ischemic, autoimmune and viral diseases, and cancer.
XX
XX Claim 4; SEQ ID NO 17; 316pp; English.
XX
XX The present invention relates to novel purified proteins (I) and their
XX coding sequences (II) (ADR89477-ADR89550), which are capable of inducing
XX apoptosis. The proteins (I) are useful as a target in screening for
XX compounds that modulate apoptosis. Compounds that modulate the expression
XX or activity of the protein may be useful in treating neurodegenerative
XX diseases, ischaemic diseases, cancer, autoimmune diseases, or viral
XX diseases. Measurement of the expression or activity of the protein may
XX also be used to diagnose or disease or a susceptibility to a disease.
XX
XX Sequence 1800 BP; 323 A; 526 C; 556 G; 395 T; 0 U; 0 Other;
XX
XX Query Match 22.6%; Score 29.2; DB 13; Length 1800;
XX Best Local Similarity 59.8%; Pred. No. 15;
XX Matches 49; Conservative 0; Mismatches 33; Indels 0; Gaps 0;
XX
XX Qy 48 ACCCGACCGAGATGGTGGGGTCTTTTGTGTGGTGTGTGACGTGTGTCCACCGT 107
XX | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
XX Db 769 ATCCCTCACCTCCTGGGGGATGGTGTGTGTGGGCTGTAACTGCTGCCCACTTC 828
XX | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
XX Qy 108 ATTATTCGGGACTAGTTCAGCG 129
XX | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
XX Db 829 ATCAATGCCTACTTGGTGGACG 850
XX | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
XX
XX RESULT 15
XX ID ABL01352 standard; DNA; 1836 BP.
XX
XX ABL01352;
XX
XX 15-MAR-2002 (first entry)
```





***This Page Blank (uspto)***

GenCore version 5.1.7  
Copyright (c) 1993 - 2006 Bioceleration Ltd.

OM nucleic - nucleic search, using sw model

Run on: March 7, 2006, 11:55:06 ; Search time 1586.98 Seconds  
(without alignments)  
3803.154 Million cell updates/sec

Title: US-10-782-899-1\_COPY\_472\_600

Perfect score: 129

Sequence: 1 tgaccttctgctgtagcga.....tattccgactgttcagcg 129

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 41078325 seqs, 23393541228 residues

Total number of hits satisfying chosen parameters: 82156650

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database :

EST:\*  
1: gb\_est1:\*  
2: gb\_est2:\*  
3: gb\_est3:\*  
4: gb\_hic:\*  
5: gb\_est4:\*  
6: gb\_est5:\*  
7: gb\_est6:\*  
8: gb\_est7:\*  
9: gb\_gss1:\*  
10: gb\_gss2:\*  
11: gb\_gss3:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

# SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
C 1	31.8	24.7	600	2	BG817831
C 2	31.4	24.3	786	8	CX773853
C 3	31.2	24.2	748	8	CV949375
C 4	31	24.0	460	7	CF927863
C 5	31	24.0	932	7	CK279022
C 6	30.8	23.9	429	3	BM522110
C 7	30.8	23.9	524	1	AA517086
C 8	30.8	23.9	938	7	CN808355
C 9	30.6	23.7	199	1	AJ282728
C 10	30.6	23.7	676	1	AV745983
C 11	30.6	23.7	790	10	CZ036908
C 12	30.2	23.4	244	7	CV609497
C 13	30.2	23.4	494	3	BM522137
C 14	30.2	23.4	528	3	BM522043
C 15	30.2	23.4	539	3	BM522233
C 16	30.2	23.4	546	2	BG817800
C 17	30.2	23.4	572	3	BM522287
C 18	30.2	23.4	577	2	BG817916
C 19	30.2	23.4	599	2	BG817905
C 20	30.2	23.4	604	7	CV614668
C 21	30.2	23.4	657	2	BG817854
C 22	30.2	23.4	682	2	BG817862

C 23	30.2	23.4	720	2	BG817630
C 24	30.2	23.4	1101	10	CNS017UC
C 25	29.8	23.1	125	6	CAB29999
C 26	29.8	23.1	486	6	CB816450
C 27	29.8	23.1	655	7	CK561781
C 28	29.8	23.1	1396	6	CD326834
C 29	29.6	22.9	187	2	BF841751
C 30	29.6	22.9	403	2	BG925693
C 31	29.6	22.9	819	7	CNI68123
C 32	29.6	22.9	903	11	CNS04GSE
C 33	29.6	22.9	937	6	CD107403
C 34	29.4	22.8	137	3	BJ305960
C 35	29.4	22.8	364	4	AK210039
C 36	29.4	22.8	437	3	BM521973
C 37	29.4	22.8	485	8	H76502
C 38	29.4	22.8	770	10	CG997493
C 39	29.4	22.8	1353	2	BI114606
C 40	29.2	22.6	231	4	AK178745
C 41	29.2	22.6	231	4	AK201694
C 42	29.2	22.6	231	4	AK208166
C 43	29.2	22.6	259	1	AI613789
C 44	29.2	22.6	282	4	AK195241
C 45	29.2	22.6	282	4	AK199429

## ALIGNMENTS

RESULT 1  
BG817831/c  
LOCUS  
DEFINITION  
ESSU0253 S.scabiei cDNA library Sarcoptes scabiei EST 22-MAY-2001  
SAS0334 5', mRNA sequence.  
600 bp mRNA linear cDNA clone  
ACCESSION  
BG817831  
VERSION  
BG817831.1 GI:14188811  
KEYWORDS  
EST.  
SOURCE  
Sarcoptes scabiei  
ORGANISM  
Sarcoptes scabiei  
Eukaryota; Metazoa; Arthropoda; Chelicerata; Arachnida; Acari;  
Acariformes; Sarcoptiformes; Astigmata; Psoroptidia; Sarcoptoides;  
Sarcoptidae; Sarcoptes.  
REFERENCE  
1 (bases 1 to 600)  
Ljunggren,B.L., Nilsson,D., Naslund,K. and Mattsson,J.G.  
Expressed sequence tag analysis of the parasitic mite Sarcoptes scabiei  
Unpublished (2001)  
Contact: Mattsson J.G.  
Department of Parasitology (SWEPAR)  
National Veterinary Institute  
SE-751 89 Uppsala, Sweden  
Tel: +46 18 674120  
Fax: +46 18  
Email: jens.mattsson@sva.se  
Similar to pit17405 scavenger recept or cysteine-rich protein precursor - sea urchin (Strongyloce nitrotus purpuratus).  
Seq primer: T3 primer  
High quality sequence stop: 600.  
FEATURES  
Location/Qualifiers  
1..600  
/organism="Sarcoptes scabiei"  
/mol\_type="mRNA"  
/db\_xref="taxon:52283"  
/clone="SAS0334"  
/note="The Sarcoptes scabiei cDNA library"  
/note="The Sarcoptes scabiei mixed lifestage library was constructed by Jens G Mattsson. cDNAs were synthesized from poly(A)+ RNA by oligo d(T) priming, size-selected and directionally cloned into the Uni-ZAP lambda vector (Stratagene). The primary library was amplified on XL1-Blue MRF' cells."

ORIGIN

Query Match

24.7%; Score 31.8; DB 2; Length 600;

```

Best Local Similarity 57.6%; Pred. No. 17;
Matches 57; Conservative 0; Mismatches 42; Indels 0; Gaps 0;

QY 2 GACCTTCTGCTAGCATTACTTCGAGCATTACTGACGACAAAGACCCGACCGAGAT 61
    |||||
Db 423 GACCATGAGGACGTCACATGTTCTCGGCATGACGACCAAGGAGGCCAATGAGAA 364
    |||||

QY 62 GGTGCGGGCTTTTGTGTGGTCTGTGACGTGTTGTC 100
    |||||
Db 363 GTTTAGGGCATCTGTGTGGTGGTCTTTTGCAATGTTTC 325

RESULT 2
CX773853/c
LOCUS CX773853.1 786 bp mRNA linear EST 25-JAN-2005
DEFINITION UI-EH-HGO-aaJ-f-23-0-UI.81 UI-EH-HGO Emiliaia huxleyi cDNA clone
ACCESSION CX773853
VERSION 1
KEYWORDS Eukaryota; Haptophyceae; Isochrysidales; Emiliaia.
SOURCE Bonaudo, M.F., Lennon, G. and Soares, M.B.
ORGANISM Normalization and subtraction: two approaches to facilitate gene
REFERENCE 1 (bases 1 to 786)
AUTHORS
TITLE Normalization and subtraction: two approaches to facilitate gene
JOURNAL Genome Res. 6 (9), 791-806 (1996)
PUBMED 8889548
COMMENT Contact: Soares, MB
Coordinated Laboratory for Computational Genomics
University of Iowa
375 Newton Road, 4156 MEBRF, Iowa City, IA 52242, USA
Tel: 319 335 8250
Fax: 319 335 9565
Email: bento-soares@uiowa.edu
Tissue Procurement: Provasoli-Guillard National Center for Culture
of Marine Phytoplankton (CCMP)
cDNA Library preparation: Dr. M. Bento Soares, University of Iowa
DNA Sequencing by: Dr. M. Bento Soares, University of Iowa
Clone Distribution: Distribution information can be found at
http://genome.uiowa.edu/distribution/dinoflagellate.html
The following repetitive elements were found in this cDNA
sequence: 148-189, >GC-rich#Low_complexity (matched complement)
Seq primer: M13 FORWARD
POLYA=Yes.

FEATURES
source Location/Qualifiers
1..786
/organism="Emiliaia huxleyi"
/mol_type="mRNA"
/strain="CCMP 371"
/db_xref="taxon:2903"
/clone="UI-EH-HGO-aaJ-f-23-0-UI"
/lab_host="DH10B (Life Technologies) (T1 phage resistant)"
/clone_lib="UI-EH-HGO"
/notes="Vector: p773-pac (Pharmacia) with a modified
polylinker; Site 1: EcoR I; Site 2: Not I; The library was
constructed according to Bonaudo, Lennon and Soares,
Genome Research, 6:791-806, 1996. First strand cDNA
synthesis was primed with an oligo-dT primer containing a
Not I site. Double stranded cDNA was ligated to an EcoR I
adaptor, digested with Not I, and cloned directionally
into p773-Pac vector. The oligonucleotide used to prime
the synthesis of first-strand cDNA contains a library tag
sequence that is located between the Not I site and the
(dT)18 tail. The sequence tag for this library is
GAAGGCTAGT. Tissue was obtained from the
Provasoli-Guillard National Center for Culture of Marine
Phytoplankton (CCMP).
TAG TISSUE=Emiliaia huxleyi coccolithophorid
TAG LIB=UI-EH-HGO
TAG_SEQ=GAAGGCTAGT"

```

## ORIGIN

```

Query Match 24.3%; Score 31.4; DB 8; Length 786;
Best Local Similarity 56.2%; Pred. No. 24;
Matches 59; Conservative 0; Mismatches 46; Indels 0; Gaps 0;

QY 4 CCTTCTGCTCGTAGCATTACTTCGAGCATTACTGACGACAAAGACCCGACCGAGATGG 63
    |||||
Db 339 CCTTCTTACGTGGCTGGTGTATCCGAGCAGCGCTCTACACCAAGACGCTACGAGCGGC 280
    |||||

QY 64 TCGGGGTCTTTTGTGTGGTGTGTGACGTGTTGCCAACCGTA 108
    |||||
Db 279 GCGCCGCGCATGACGCTCTCGAGGAGCGACGTGATGCAATCCGCA 235
    |||||

```

## RESULT 3

```

CV949375/c
LOCUS CV949375.1 748 bp mRNA linear EST 25-JAN-2005
DEFINITION PVrpb13369 zoospores, purified Phytophthora infestans cDNA, mRNA
sequence.
ACCESSION CV949375
VERSION 1
KEYWORDS Phytophthora infestans (potato late blight agent)
SOURCE Phytophthora infestans
ORGANISM Eukaryota; stramenopiles; Oomycetes; Pythiales; Pythiaceae;
Phytophthora.
REFERENCE 1 (bases 1 to 748)
AUTHORS Randall, T., Dwyer, R.A., Huitema, E., Beyer, K., Cvitanich, C.,
Kelkar, H., Fong, A.M., Gates, K., Roberts, S., Yatskan, E., Gaffney, T.,
Law, M., Testa, A., Torto-Alalibo, A., Zhang, M., Zheng, L., Mueller, E.,
Windsass, J., Binder, A., Birch, P.R.J., Gisi, U., Govers, F., Gow, N.A.,
Mauch, F., van West, P., Waugh, M.E., Yu, J., Boller, T., Kamoun, S.,
Lam, S.T. and Judelson, H.S.
TITLE Large-scale gene discovery in the oomycete Phytophthora infestans
reveals likely components of phytopathogenicity shared with true
fungi
JOURNAL Mol. Plant-Microbe Interact. 18 (3), 229-243 (2005)
PUBMED 15782637
COMMENT Contact: Judelson HS
Department of Plant Pathology
University of California
Webber Hall, Riverside, CA 92521, USA
Tel: 909 787 4199
Fax: 909 787 4294
Email: howard.judelson@ucr.edu.
Location/Qualifiers
1..748
/organism="Phytophthora infestans"
/mol_type="mRNA"
/strain="88069"
/db_xref="taxon:4787"
/sex="A1"
/clone_lib="zoospores, purified"
/notes="Vector: pSPORT1"

```

## FEATURES

```

source Location/Qualifiers
1..748
/organism="Phytophthora infestans"
/mol_type="mRNA"
/strain="88069"
/db_xref="taxon:4787"
/sex="A1"
/clone_lib="zoospores, purified"
/notes="Vector: pSPORT1"

```

## ORIGIN

```

Query Match 24.2%; Score 31.2; DB 8; Length 748;
Best Local Similarity 53.2%; Pred. No. 27;
Matches 66; Conservative 0; Mismatches 58; Indels 0; Gaps 0;

QY 2 GACCTTCTGCTCGTAGCGATTACTTCGAGCATTACTGACGACAAAGACCCGACCGAGAT 61
    |||||
Db 145 GAACCTGACGCTCTCGGACGACGCGTCGCGCTTGACGACGCTACATCGTTGGATTGAGCT 86
    |||||

QY 62 GGTGCGGGCTTTTGTGTGGTGTGTGACGTGTTGCCAACCGTATTATTCGGGACTA 121
    |||||
Db 85 GGTGCGGTAACTGCTCTCGAGGTGGACGAGCGGTGGTGGTCCGCCGGAATTCGGGACCG 26
    |||||

QY 122 GTTC 125
    |||
Db 25 GTAC 22
    |||

```

```

RESULT 4
CF927863
LOCUS
DEFINITION
ACCESSION
VERSION
KEYWORDS
SOURCE
ORGANISM

CF927863
1af58f02.y1 SlEP Mus musculus cDNA 5' similar to TR:Q9UJZ5 Q9UJZ5
PRESENTINLIN-ASSOCIATED PROTEIN. 7, mRNA sequence.
CF927863
CF927863.1 GI:38274486
EST.
Mus musculus (house mouse)
Mus musculus
Eukaryota; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Euarchontoglires; Glires; Rodentia;
Sciurognathi; Muridae; Murinae; Mus.
1 (bases 1 to 460)
Wyllie, T., Theising, B., Bowers, Y., Gibbons, M., Ritter, E., Bannet, J.,
Runko, I., Tsagareishvili, R., Belaygorod, L., Grow, A., Meguire, L.,
Waterston, R. and Wilson, R.
WashU Stem Cell EST Project
Unpublished (2002)
Contact: Jeff Gordon and Mike Lovett
WashU Human Genetics Division
Washington University School of Medicine
Library materials provided by: Stappenbeck Th.S. and Gordon J.I.
Library constructed by: Korshunova Y. and Lovett M. DNA sequencing
by: Washington University Genome Sequencing Center For information
on obtaining a clone please contact: Rose Tidwell
(rtidwell@genetics.wustl.edu)
Seq primer: -40RP from Gibco
High quality sequence stop: 460.
Location/Qualifiers
FEATURES
source
1..460
/organism="Mus musculus"
/mol_type="mRNA"
/db_xref="taxon:10090"
/tissue_type="small intestinal epithelial progenitors"
/lab_host="DH5alpha Ultra Max cells (Invitrogen)"
/clone_lib="GIEP"
/notes="Vector: pAMP1; The library was synthesized with
modified SMART primers with dUTP at the end. After
treatment with UDG the cDNA was cloned in pAMP1 vector by
annealing."
ORIGIN
Query Match 24.0%; Score 31; DB 7; Length 460;
Best Local Similarity 59.8%; Pred. No. 29;
Matches 52; Conservative 0; Mismatches 35; Indels 0; Gaps 0;

QY 43 CAAAGACCCGACGAGATGTCGGGCTCTTTTGTGTGCTGTGACGTGTGTCCCA 102
Db ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
7 CTATTACCTCACTAAAGGGGGGATGCTGTTTCTTGTGGGCTGTAACTGCTGGCCC 66

QY 103 ACCGTATATTCGGACTAGTTACGC 129
Db ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||

Db 67 ACTTCATCAATGCCCTACTTGGTGACG 93

RESULT 5
CK279022/c
LOCUS
DEFINITION
ACCESSION
VERSION
KEYWORDS
SOURCE
ORGANISM

CK279022
EST725100 potato abiotic stress cDNA library Solanum tuberosum cDNA
clone POAE963 5' end, mRNA sequence.
CK279022
CK279022.1 GI:39836000
EST.
Solanum tuberosum (potato)
Solanum tuberosum
Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
Spermatophyta; Magnoliophyta; eudicotyledons; core eudicotyledons;
asterids; lamids; Solanales; Solanaceae; Solanum.
1 (bases 1 to 932)
Buell, C.R., Hart, A., Zismann, V., Karamycheva, S.A. and Baker, B.
REFERENCE
AUTHORS

TITLE
JOURNAL
COMMENT

Generation of ESTs from abiotic stressed potato tissue
Unpublished (2003)
Contact: Robin Buell
The Institute for Genomic Research
9712 Medical Center Dr, Rockville, MD 20850, USA
Email: potato-array@tigr.org
Clones can be requested from the University of Arizona Genomics
Institute via http://genome.arizona.edu/orders/
Seq primer: ATT TAG GTG ACA CTA TAG.
Location/Qualifiers
FEATURES
source
1..932
/organism="Solanum tuberosum"
/mol_type="mRNA"
/cultivar="Kennebec"
/db_xref="taxon:4113"
/clone="POAE963"
/tissue_type="abiotic stress treated leaf and root tissue"
/lab_host="DH10B-Tona"
/clone_lib="potato abiotic stress cDNA library"
/notes="Vector: pCMVSPORT6.1; Site 1: EORI; Site 2: NotI;
supplier: Solanum tuberosum var. Kennebec plants were
grown from cuttings on a 16hr light/8 hr dark cycle at 25
C for 3-4 weeks. Abiotic stress conditions were applied to
four separate sets of plants. Set 1 involved saturation of
the soil with 150 mM NaCl and tissues were harvested at
following application of the salt stress (leaves: 2hr,
6hr, 12hr, 1d, 2d, and 4d; roots: 2hr, 6hr, 12hr, and 2d).
Set 2 were grown under the standard conditions and then
were water stressed by withdrawal of further watering
applications. Drought stressed plants were harvested after
cessation of watering (leaves: 3d, 5d, and 7d; roots: 3d
and 5d). Set 3 were grown under the standard conditions
and then were cold stressed by placement at 4 C. Cold
stressed leaves were harvested at 2 hr, 6 hr, 12 hr, 1 d,
and 4d and roots were harvested at 2 hr, 6 hr, 12 hr, 1 d,
2d. Set 4 were grown under the standard conditions and
then were heat stressed by placement at 35 C. Heat
stressed leaves were harvested at 2 hr, 6 hr, 12 hr, 1 d,
2d and 4d and heat-stressed roots were harvested at 6 hr,
12 hr, 1 d, and 4d. RNA was isolated from all tissues and
equal RNA from each tissue and stress was pooled to
construct the cDNA library. RNA sample."
ORIGIN
Query Match 24.0%; Score 31; DB 7; Length 932;
Best Local Similarity 57.9%; Pred. No. 33;
Matches 55; Conservative 0; Mismatches 40; Indels 0; Gaps 0;

QY 5 CTTCTGCTCGTACGATTACTTCGAGCATTAACGACAAAGACCCGACGATGGT 64
Db ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
844 CTTCTGTTAATATTGAATAGAAAGCTGATTTGAGGAGAAAGAGAGTGTGATGAT 785

QY 65 CGGGGCTCTTTTCTGTGGTGTGACGTGTGT 99
Db ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
784 GTTGTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTT 750

RESULT 6
BM522110/c
LOCUS
DEFINITION
ACCESSION
VERSION
KEYWORDS
SOURCE
ORGANISM

BM522110
ESSU0647 S.scabiei cDNA library Sarcopetes scabiei cDNA clone
SAS0777 5', mRNA sequence.
BM522110
BM522110.1 GI:18706032
EST.
Sarcopetes scabiei
Sarcopetes scabiei
Eukaryota; Metazoa; Arthropoda; Chelicerata; Arachnida; Acari;
Acariformes; Sarcopetiformes; Astigmata; Psoroptidia; Sarcopetidae;
Sarcopetes.
1 (bases 1 to 429)
Ljunggren, E.L., Nilsson, D., Naslund, K. and Mattsson, J.G.
Expressed sequence tag analysis of the parasitic mite Sarcopetes
TITLE

```



```

Query Match      23.9%; Score 30.8; DB 7; Length 938;
Best Local Similarity 63.5%; Pred. No. 38;
Matches 47; Conservative 0; Mismatches 27; Indels 0; Gaps 0;

QY 7 TCCTGCTGATGACGATTTCTGAGCATTTCTGACGACAAAGACCCGACCGAGATGCTCG 66
Db 330 TTTGGGCGGGGGGCTCTTCGCGCTTCTGACGACAAAGTCGACGCGCGCTTTGGAGT 271

QY 67 GGGTCTTTTGGTTG 80
Db 270 AGATCTTGGGGTTG 257

RESULT 9
AJ282728/c
LOCUS      199 bp      mRNA      linear      EST 30-JUN-2000
DEFINITION 4A3A-ABB-D-05-R Anopheles gambiae immune competent 4A3A Anopheles
SOURCE      gambiae cDNA clone 4A3A-ABB-D-05, mRNA sequence.
ACCESSION   AJ282728
VERSION     1
KEYWORDS    EST.
ORGANISM    Anopheles gambiae (African malaria mosquito)
REFERENCE   1 (bases 1 to 199)
AUTHORS     Dimopoulos, G., Casavant, T.L., Chang, S., Scheetz, T., Roberts, C.,
            Donohue, M., Schultz, J., Benes, V., Bork, P., Ansoerge, W., Soares, M.B.
            and Kafatos, F.C.
TITLE       Anopheles gambiae pilot gene discovery project: identification of
            mosquito innate immunity genes from expressed sequence tags
            generated from immune-competent cell lines
JOURNAL     Proc. Natl. Acad. Sci. U.S.A. 97 (12), 6619-6624 (2000)
PUBMED     10841561
COMMENT     Contact: Dimopoulos G
            Fotis C. Kafatos Laboratory
            European Molecular Biology Laboratory
            Meyerhofstrasse 1, 69117 Heidelberg, Germany.
FEATURES    source
            1..199
            /organism="Anopheles gambiae"
            /mol_type="mRNA"
            /strain="4A x/r"
            /db_xref="taxon:7165"
            /clone="4A3A-ABB-D-05"
            /cell_line="immune competent 4A3A"
            /lab_host="E. coli DH10B"
            /notes="Vector: p773D-Pac (Pharmacia) with a modified
            polylinker; Site 1: EcoRI; Site 2: NotI; sequenced from
            forward priming site which reads from the 3' end of the
            cDNA. The 4A3A is a directionally cloned and normalized
            cDNA library that was constructed from the 4A3A cell line
            oligo-T primed cDNA according to: Bonaldo, Lennon & Soares
            (1996) : Normalization and Subtraction: Two approaches To
            Facilitate Gene Discovery, Genome Research 6, 791-806."

ORIGIN
Query Match      23.7%; Score 30.6; DB 1; Length 199;
Best Local Similarity 55.0%; Pred. No. 34;
Matches 60; Conservative 0; Mismatches 49; Indels 0; Gaps 0;

QY 12 TCCTAGGATTTCTGAGCATTTCTGACGACAAAGACCCGACCGAGATGTCGGGTC 71
Db 184 TGGGAGCAATTTAAACGGCGCACTACTAACGGTAACGGACCAACACGTAATGTCCCGGGT 125

QY 72 TTTTGTGTGGTCTGAGCGTTGTCCTCAACCGTATTATTCGGACT 120
Db 124 CTTTGTGTGTTCATTTCGGCGGTAGTAAATCGGCTTCTACTAGACT 76

Query Match      23.7%; Score 30.6; DB 1; Length 676;
Best Local Similarity 53.8%; Pred. No. 42;
Matches 63; Conservative 0; Mismatches 54; Indels 0; Gaps 0;

QY 2 GACCTTCTGCTCTAGCGATTACTTTCGAGCATTTCTGACGACAAAGACCCGACCGAGAT 61
Db 435 GACCGCTCTCTGGATCTCCCGCGCGAGCCTTCTCAGAGACATCGAGCGGTCTGAGAG 494

QY 62 GGTGGGGCTTTTCTTGTGGTGTGTCGACGTTGTCCACCGTATTATTCGGA 118
Db 495 CCTCTGGGACGCTTTGTGTGTGTGTAACTCAACCTTAAGATAATGGA 551

NOTE=Vector: pBluescript sk(-); Site_1: EcoRI; Site_2:
XhoI"

RESULT 11
CZ036908/c
LOCUS      790 bp      DNA      linear      GSS 10-JAN-2005
DEFINITION OM_Ba0016122.f OM_Ba Oryza minuta genomic clone OM_Ba0016122 5',
SOURCE      genomic survey sequence.
ACCESSION   CZ036908
VERSION     1
KEYWORDS    GSS.
ORGANISM    Oryza minuta
            Oryza minuta
            Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
            Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;
            Ehrhartoideae; Oryzaceae; Oryza.
            1 (bases 1 to 790)
REFERENCE   1
AUTHORS     Kim, H., Collura, K., Wisseotski, M., Byrne, M., Stum, D., Smart, D.,
            Rao, K., Luo, M., Jetty, R., Kudrna, D., Muller, C., Soderlund, C. and

```

Wing, R.  
 OMAP (Oryza Map Alignment Project) - Arizona Genomics Institute  
 Unpublished (2005)  
 Contact: Rod A. Wing  
 Arizona Genomics Institute  
 University of Arizona  
 Forbes Building Room 303, Tucson, AZ 85721-0036, USA  
 Tel: 520 626 9595  
 Fax: 520 621 1259  
 Email: rwing@genome.arizona.edu  
 Plate: 0016 row: I column: 22  
 Class: BAC ends.

**FEATURES**  
 source  
 1..790  
 /organism="Oryza minuta"  
 /mol\_type="genomic DNA"  
 /db\_xref="taxon:63629"  
 /clone="OM\_Ba0016122"  
 /tissue\_type="leaves"  
 /lab\_host="DH108"  
 /clone\_lib="OM\_Ba"  
 /notes="Vector: pCUGIBAC1; Site\_1: HindIII; Site\_2: HindIII"

**ORIGIN**  
 Query Match 23.7%; Score 30.6; DB 10; Length 790;  
 Best Local Similarity 60.0%; Pred. No. 43; Mismatches 0; Indels 0; Gaps 0;  
 Matches 51; Conservative 0;

Qy 42 ACAAGACCCCGACGAGATGTCGGGGTCTTTTGTGTGGTCTGTGACGTGTGTCC 101  
 |||||  
 Db 510 ACGACGAGACCATCGAATGTCGGTAGTTTTTTGGCTTCGCTACGTGCGGTAC 451  
 |||||

Qy 102 AACCCTATTATTCGGACTAGTTCA 126  
 |||||  
 Db 450 AACTGTATAAATAAGTGGAAATTAA 426  
 |||||

**RESULT 12**  
 CV609497/c  
 LOCUS zcm31a01.y2\_F\_HC7CDNA\_Ajellomyces capsulatus CDNA 5', mRNA  
 DEFINITION sequence.  
 ACCESSION CV609497  
 VERSION CV609497.1 GI:54597079  
 KEYWORDS EST.  
 SOURCE Ajellomyces capsulatus  
 ORGANISM Ajellomyces capsulatus  
 Eukaryota; Fungi; Ascomycota; Pezizomycotina; Eurotiomycetes;  
 Onygenales; Ajellomycetaceae; Ajellomyces.  
 1 (bases 1 to 244)  
 /clone\_lib="S. scabiei cDNA library"  
 /note="The Sarcptes scabiei mixed life stage library was constructed by Jens G Mattsson. cDNAs were synthesized from poly(A)+ RNA by oligo d(T) priming, size-selected and directionally cloned into the Uni-ZAP lambda vector (Stratagene). The primary library was amplified on XL1-Blue MRF' cells."

**REFERENCE**  
 AUTHORS Magrini, V., Sil, A., Goldman, W. and Mardis, R.  
 TITLE Genomic Resources for Histoplasma Sequencing  
 HISTOPLASMA CAPSULATUM SEQUENCING PROJECT  
 UNPUBLISHED (2004)  
 CONTACT: Vincent Magrini  
 Mardis/Goldman/Sil Genomic Resources for Histoplasma Sequencing  
 Washington University School of Medicine  
 4444 Forest Park Parkway, Box 8501, St. Louis, MO 63108, USA  
 Tel: 314 286 1800  
 Fax: 314 286 1810  
 Email: vmagrini@wustl.edu  
 Contact Vincent Magrini (vmagrini@wustl.edu) for further information relating to organism, libraries, or clone availability.  
 Seq primer: H7CDNA-F2.  
 Location/Qualifiers  
 1..244  
 /organism="Ajellomyces capsulatus"  
 /mol\_type="mRNA"  
 /strain="G217B"  
 /db\_xref="taxon:5037"  
 /clone\_lib="F\_HC7CDNA"  
 /note="Vector: pAS209; Site\_1: NotI; Site\_2: SalI;

**FEATURES**  
 source  
 1..244  
 /organism="Ajellomyces capsulatus"  
 /mol\_type="mRNA"  
 /strain="G217B"  
 /db\_xref="taxon:5037"  
 /clone\_lib="F\_HC7CDNA"  
 /note="Vector: pAS209; Site\_1: NotI; Site\_2: SalI;

**ORIGIN**  
 Query Match 23.4%; Score 30.2; DB 7; Length 244;  
 Best Local Similarity 62.7%; Pred. No. 48; Mismatches 0; Indels 0; Gaps 0;  
 Matches 47; Conservative 0;

Qy 13 CGTAGCGATTACTTCGAGCATTTACTACGACGACAAAGACCCGACCGAGATGGTCGGGTCT 72  
 |||||  
 Db 233 CGTAGAGATGACGTCCTCTTTATTGAGGGCATAGACACATCCAGCGATGTGTCGGTCT 174  
 |||||

Qy 73 TTTTGTGTGGTGCT 87  
 |||||  
 Db 173 TTCTCTTGTGCTGCT 159  
 |||||

**RESULT 13**  
 BM522137/c  
 LOCUS ESSU0674 S.scabiei cDNA library Sarcptes scabiei cDNA clone  
 DEFINITION SAS0807 5', mRNA sequence.  
 ACCESSION BM522137  
 VERSION BM522137.1 GI:18706191  
 KEYWORDS EST.  
 SOURCE Sarcptes scabiei  
 ORGANISM Sarcptes scabiei  
 Eukaryota; Metazoa; Arthropoda; Chelicerata; Arachnida; Acari;  
 Acariformes; Sarcptiformes; Astigmata; Peoroptidia; Sarcptoidae;  
 Sarcptidae; Sarcptes.  
 1 (bases 1 to 494)  
 /clone\_lib="S.scabiei cDNA library"  
 /note="The Sarcptes scabiei mixed life stage library was constructed by Jens G Mattsson. cDNAs were synthesized from poly(A)+ RNA by oligo d(T) priming, size-selected and directionally cloned into the Uni-ZAP lambda vector (Stratagene). The primary library was amplified on XL1-Blue MRF' cells."

**REFERENCE**  
 AUTHORS Ljunggren, E.L., Nilsson, D., Naglund, K. and Mattsson, J.G.  
 TITLE Expressed sequence tag analysis of the parasitic mite Sarcptes scabiei  
 JOURNAL Unpublished (2001)  
 COMMENT Contact: Mattsson J.G.  
 Department of Parasitology (SWEPAR)  
 National Veterinary Institute  
 SE-751 89 Uppsala, Sweden  
 Tel: +46 18 674120  
 Fax: +46 18

**FEATURES**  
 source  
 1..494  
 /organism="Sarcptes scabiei"  
 /mol\_type="mRNA"  
 /db\_xref="taxon:52283"  
 /clone="SAS0807"  
 /clone\_lib="S.scabiei cDNA library"  
 /note="The Sarcptes scabiei mixed life stage library was constructed by Jens G Mattsson. cDNAs were synthesized from poly(A)+ RNA by oligo d(T) priming, size-selected and directionally cloned into the Uni-ZAP lambda vector (Stratagene). The primary library was amplified on XL1-Blue MRF' cells."

**ORIGIN**  
 Query Match 23.4%; Score 30.2; DB 3; Length 494;  
 Best Local Similarity 56.6%; Pred. No. 54; Mismatches 0; Indels 0; Gaps 0;  
 Matches 56; Conservative 0;

Qy 2 GACCTTCTGCTCGTAGCGATTACTTCGAGCATTTACTACGACGACAAAGACCCGACCGAGAT 61  
 |||||  
 Db 135 GACCATGAGACGTGCACATGTTCTCGGGCATGAACGACCAAGGAGGCCCAATGAGAA 76  
 |||||

Qy 62 GGTCCGGGTCTTTTGTGTGTGCTGTGACGTGTGTC 100  
 |||||  
 Db 75 GTTTAGGGCATCTGTGTGTGAGGCTTTTTCATGTCTTC 37  
 |||||



RESULT 14  
BM522043/c  
LOCUS  
DEFINITION  
ESSU0580 S.scabiei cDNA library Sarcopes scabiei cDNA clone  
SAS0706 5', mRNA sequence.  
ACCESSION  
VERSION  
KEYWORDS  
SOURCE  
EST.  
ORGANISM  
Sarcopes scabiei  
Eukaryota; Metazoa; Arthropoda; Chelicerata; Arachnida; Acari;  
Acariiformes; Sarcotiformes; Astigmata; Psoroptidia; Sarcotoides;  
Sarcotidae; Sarcopes.  
REFERENCE  
AUTHORS  
TITLE  
Expressed sequence tag analysis of the parasitic mite Sarcopes  
scabiei

JOURNAL  
COMMENT  
Unpublished (2001)  
Contact: Mattsson J.G.  
Department of Parasitology (SWEPAR)  
National Veterinary Institute  
SE-751 89 Uppsala, Sweden  
Tel: +46 18 674120  
Fax: +46 18

Similar to gi|7512096|pir|T17405 scavenger receptor cysteine-rich  
protein precursor - sea urchin [Strongylocentrotus purpuratus]  
Seq primer: T3 primer  
High quality sequence stop: 528.

FEATURES  
source  
Location/Qualifiers  
1..528  
/organism="Sarcopes scabiei"  
/mol\_type="mRNA"  
/db\_xref="taxon:52283"  
/clone\_lib="SAS0706"  
/notes="The Sarcopes scabiei mixed life stage library was  
constructed by Jens G Mattsson. cDNAs were synthesized  
from poly(A)+ RNA by oligo d(T) priming, size-selected and  
directionally cloned into the Uni-ZAP lambda vector  
(Stratagene). The primary library was amplified on  
XL1-Blue MRF' cells."

ORIGIN  
Query Match 23.4%; Score 30.2; DB 3; Length 528;  
Best Local Similarity 56.6%; Pred. No. 55;  
Matches 56; Conservative 0; Mismatches 43; Indels 0; Gaps 0;  
QY 2 GACCTTCTGCTAGCGATTACTTCGAGCATTACTGACGACAAAGACCCGACCGAGAT 61  
Db 387 GACCATGAGGACGTGCACATGTTCTCGGCATGACGACCAAGGAGGCCAATGAGAT 328  
QY 62 GGTGCGGGTCTTTTGTGTGGTCTGTGACGTGTGTC 100  
Db 327 GTTTAGGGCATCTGTGTGAGGTCTTTTGCAATGTCTTC 289

Search completed: March 7, 2006, 13:19:43  
Job time : 1595.98 secs

RESULT 15  
BM522233/c  
LOCUS  
DEFINITION  
ESSU0771 S.scabiei cDNA library Sarcopes scabiei cDNA clone  
SAS0908 5', mRNA sequence.  
ACCESSION  
VERSION  
KEYWORDS  
SOURCE  
EST.  
ORGANISM  
Sarcopes scabiei  
Eukaryota; Metazoa; Arthropoda; Chelicerata; Arachnida; Acari;  
Acariiformes; Sarcotiformes; Astigmata; Psoroptidia; Sarcotoides;  
Sarcotidae; Sarcopes.  
REFERENCE  
AUTHORS  
TITLE  
Expressed sequence tag analysis of the parasitic mite Sarcopes  
scabiei

JOURNAL  
COMMENT  
Unpublished (2001)  
Contact: Mattsson J.G.  
Department of Parasitology (SWEPAR)  
National Veterinary Institute  
SE-751 89 Uppsala, Sweden  
Tel: +46 18 674120  
Fax: +46 18

Similar to gi|7512096|pir|T17405 scavenger receptor cysteine-rich  
protein precursor - sea urchin [Strongylocentrotus purpuratus]  
Seq primer: T3 primer  
High quality sequence stop: 528.

FEATURES  
source  
Location/Qualifiers  
1..528  
/organism="Sarcopes scabiei"  
/mol\_type="mRNA"  
/db\_xref="taxon:52283"  
/clone\_lib="SAS0706"  
/notes="The Sarcopes scabiei mixed life stage library was  
constructed by Jens G Mattsson. cDNAs were synthesized  
from poly(A)+ RNA by oligo d(T) priming, size-selected and  
directionally cloned into the Uni-ZAP lambda vector  
(Stratagene). The primary library was amplified on  
XL1-Blue MRF' cells."

ORIGIN  
Query Match 23.4%; Score 30.2; DB 3; Length 528;  
Best Local Similarity 56.6%; Pred. No. 55;  
Matches 56; Conservative 0; Mismatches 43; Indels 0; Gaps 0;  
QY 2 GACCTTCTGCTAGCGATTACTTCGAGCATTACTGACGACAAAGACCCGACCGAGAT 61  
Db 387 GACCATGAGGACGTGCACATGTTCTCGGCATGACGACCAAGGAGGCCAATGAGAT 328  
QY 62 GGTGCGGGTCTTTTGTGTGGTCTGTGACGTGTGTC 100  
Db 327 GTTTAGGGCATCTGTGTGAGGTCTTTTGCAATGTCTTC 289

Search completed: March 7, 2006, 13:19:43  
Job time : 1595.98 secs

scabiei  
Unpublished (2001)  
Contact: Mattsson J.G.  
Department of Parasitology (SWEPAR)  
National Veterinary Institute  
SE-751 89 Uppsala, Sweden  
Tel: +46 18 674120  
Fax: +46 18

Similar to gi|7512096|pir|T17405 scavenger receptor cysteine-rich  
protein precursor - sea urchin [Strongylocentrotus purpuratus]  
Seq primer: T3 primer  
High quality sequence stop: 539.

FEATURES  
source  
Location/Qualifiers  
1..539  
/organism="Sarcopes scabiei"  
/mol\_type="mRNA"  
/db\_xref="taxon:52283"  
/clone\_lib="SAS0908"  
/notes="The Sarcopes scabiei mixed life stage library was  
constructed by Jens G Mattsson. cDNAs were synthesized  
from poly(A)+ RNA by oligo d(T) priming, size-selected and  
directionally cloned into the Uni-ZAP lambda vector  
(Stratagene). The primary library was amplified on  
XL1-Blue MRF' cells."

ORIGIN  
Query Match 23.4%; Score 30.2; DB 3; Length 539;  
Best Local Similarity 56.6%; Pred. No. 55;  
Matches 56; Conservative 0; Mismatches 43; Indels 0; Gaps 0;  
QY 2 GACCTTCTGCTAGCGATTACTTCGAGCATTACTGACGACAAAGACCCGACCGAGAT 61  
Db 402 GACCATGAGGACGTGCACATGTTCTCGGCATGACGACCAAGGAGGCCAATGAGAT 343  
QY 62 GGTGCGGGTCTTTTGTGTGGTCTGTGACGTGTGTC 100  
Db 342 GTTTAGGGCATCTGTGTGAGGTCTTTTGCAATGTCTTC 304

Search completed: March 7, 2006, 13:19:43  
Job time : 1595.98 secs

***This Page Blank (uspto)***

Result No.	Score	Query			DB	ID	Description
		Match	Length	Time			
1	29.4	22.8	870	3	US-09-583-110-1324	Sequence 1324, Ap	
2	29.4	22.8	885	3	US-09-107-433-306	Sequence 306, App	
3	29.4	22.8	194915	3	US-09-949-016-15584	Sequence 15584, A	
4	29	22.5	601	3	US-09-949-016-68059	Sequence 68059, A	
5	29	22.5	264665	3	US-09-949-016-13747	Sequence 13747, A	
C	6	28.6	22.2	1146	3	US-09-252-991A-2602	Sequence 2602, Ap
	7	28.6	22.2	2184	3	US-09-252-991A-2341	Sequence 2241, Ap
8	28.6	22.2	2858	3	US-09-252-991A-2311	Sequence 2311, Ap	
9	28.4	22.0	936	2	US-08-602-359A-31	Sequence 31, Appl	
C	10	27.4	21.2	1160	3	US-09-270-767-13705	Sequence 13705, A
11	27.4	21.2	1951	3	US-10-144-929-56	Sequence 56, Appl	
C	12	26.6	20.6	601	3	US-09-949-016-125432	Sequence 125432
13	26.6	20.6	139552	3	US-09-949-016-15300	Sequence 15300, A	
C	14	26.4	20.5	14707	3	US-09-312-762A-3	Sequence 3, Appli
C	15	26.2	20.3	601	3	US-09-949-016-143314	Sequence 143314
C	16	26.2	20.3	601	3	US-09-949-016-143485	Sequence 143485
C	17	26.2	20.3	746	3	US-09-380-418C-1	Sequence 1, Appli
C	18	26.2	20.3	907	3	US-08-672-850-7	Sequence 7, Appli
C	19	26.2	20.3	907	3	US-08-672-850-11	Sequence 11, Appli
C	20	26.2	20.3	907	3	US-09-565-177A-7	Sequence 7, Appli
C	21	26.2	20.3	907	3	US-09-565-177A-11	Sequence 11, Appli
C	22	26.2	20.3	1001	3	US-08-672-850-10	Sequence 10, Appli
C	23	26.2	20.3	1001	3	US-09-565-177A-10	Sequence 10, Appli
C	24	26.2	20.3	85912	3	US-09-949-016-12362	Sequence 12362, A



; CURRENT APPLICATION NUMBER: US/09/949,016  
; CURRENT FILING DATE: 2000-04-14  
; PRIOR APPLICATION NUMBER: 60/241,755  
; PRIOR FILING DATE: 2000-10-20  
; PRIOR APPLICATION NUMBER: 60/237,768  
; PRIOR FILING DATE: 2000-10-03  
; PRIOR APPLICATION NUMBER: 60/231,498  
; PRIOR FILING DATE: 2000-09-08  
; NUMBER OF SEQ ID NOS: 207012  
; SOFTWARE: FastSeq for Windows Version 4.0  
; SEQ ID NO 13747  
; LENGTH: 264665  
; TYPE: DNA  
; ORGANISM: Human  
; US-09-949-016-13747

Query Match 22.5%; Score 29; DB 3; Length 264665;  
Best Local Similarity 58.8%; Pred. No. 17;  
Matches 50; Conservative 0; Mismatches 35; Indels 0; Gaps 0;  
QY 23 ACTTCGAGCATTACTGACGCAAGACCCCGACCGAGATGTCGGGTCCTTTTGTGTG 82  
Db 52981 AGTGCTAGGATTACAGCGGTGAGCCACCGTGC CGGCAGACCTGGCTCTTTCTTTGTT 53040  
QY 83 GTGCTGTGACGTGTTTCACACCGT 107  
Db 53041 TTTTTCGACGAGTGTCCCTCTGT 53065

RESULT 6  
US-09-252-991A-2602/c  
; Sequence 2602, Application US/09252991A  
; Patent No. 6551795  
; GENERAL INFORMATION:  
; APPLICANT: Marc J. Rubenfield et al.  
; TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO PSEUDOMONAS  
; FILE REFERENCE: 107196.136  
; CURRENT APPLICATION NUMBER: US/09/252,991A  
; PRIOR FILING DATE: 1999-02-18  
; PRIOR APPLICATION NUMBER: US 60/074,788  
; PRIOR FILING DATE: 1998-02-18  
; PRIOR APPLICATION NUMBER: US 60/094,190  
; PRIOR FILING DATE: 1998-07-27  
; NUMBER OF SEQ ID NOS: 33142  
; SEQ ID NO 2602  
; LENGTH: 1146  
; TYPE: DNA  
; ORGANISM: Pseudomonas aeruginosa  
; US-09-252-991A-2602

Query Match 22.2%; Score 28.6; DB 3; Length 1146;  
Best Local Similarity 57.1%; Pred. No. 2.6;  
Matches 52; Conservative 0; Mismatches 39; Indels 0; Gaps 0;  
QY 39 ACACAAAGACCCCGACCGAGATGTCGGGTCCTTTTGTGTGCTGTGACGTGTTG 98  
Db 906 ACATCAAGACCTGCGCGAACTCGGGGTGATCTTCTGATGTTCTGCCTCGGCCTGGAG 847  
QY 99 TCCAAACCGTATTATTCGGGACTAGTTCAGCG 129  
Db 846 TTCAGCCTGGCAAGCTCTTCCAGGTCGGCG 816

RESULT 7  
US-09-252-991A-2241  
; Sequence 2241, Application US/09252991A  
; Patent No. 6551795  
; GENERAL INFORMATION:  
; APPLICANT: Marc J. Rubenfield et al.  
; TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO PSEUDOMONAS  
; FILE REFERENCE: 107196.136

; CURRENT APPLICATION NUMBER: US/09/252,991A  
; CURRENT FILING DATE: 1999-02-18  
; PRIOR APPLICATION NUMBER: US 60/074,788  
; PRIOR FILING DATE: 1998-02-18  
; PRIOR APPLICATION NUMBER: US 60/094,190  
; PRIOR FILING DATE: 1998-07-27  
; NUMBER OF SEQ ID NOS: 33142  
; SEQ ID NO 2241  
; LENGTH: 2184  
; TYPE: DNA  
; ORGANISM: Pseudomonas aeruginosa  
; US-09-252-991A-2241

Query Match 22.2%; Score 28.6; DB 3; Length 2184;  
Best Local Similarity 57.1%; Pred. No. 3.4;  
Matches 52; Conservative 0; Mismatches 39; Indels 0; Gaps 0;  
QY 39 ACACAAAGACCCCGACCGAGATGTCGGGTCCTTTTGTGTGCTGTGACGTGTTG 98  
Db 1538 ACATCAAGACCTGCGCGAACTCGGGGTGATCTTCTGATGTTCTGCCTCGGCCTGGAG 1597  
QY 99 TCCAAACCGTATTATTCGGGACTAGTTCAGCG 129  
Db 1598 TTCAGCCTGGCAAGCTCTTCCAGGTCGGCG 1628

RESULT 8  
US-09-252-991A-2311  
; Sequence 2311, Application US/09252991A  
; Patent No. 6551795  
; GENERAL INFORMATION:  
; APPLICANT: Marc J. Rubenfield et al.  
; TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO PSEUDOMONAS  
; FILE REFERENCE: 107196.136  
; CURRENT APPLICATION NUMBER: US/09/252,991A  
; CURRENT FILING DATE: 1999-02-18  
; PRIOR APPLICATION NUMBER: US 60/074,788  
; PRIOR FILING DATE: 1998-02-18  
; PRIOR APPLICATION NUMBER: US 60/094,190  
; PRIOR FILING DATE: 1998-07-27  
; NUMBER OF SEQ ID NOS: 33142  
; SEQ ID NO 2311  
; LENGTH: 2868  
; TYPE: DNA  
; ORGANISM: Pseudomonas aeruginosa  
; US-09-252-991A-2311

Query Match 22.2%; Score 28.6; DB 3; Length 2868;  
Best Local Similarity 57.1%; Pred. No. 3.8;  
Matches 52; Conservative 0; Mismatches 39; Indels 0; Gaps 0;  
QY 39 ACACAAAGACCCCGACCGAGATGTCGGGTCCTTTTGTGTGCTGTGACGTGTTG 98  
Db 1177 ACATCAAGACCTGCGCGAACTCGGGGTGATCTTCTGATGTTCTGCCTCGGCCTGGAG 1236  
QY 99 TCCAAACCGTATTATTCGGGACTAGTTCAGCG 129  
Db 1237 TTCAGCCTGGCAAGCTCTTCCAGGTCGGCG 1267

RESULT 9  
US-08-602-359A-31  
; Sequence 31, Application US/08602359A  
; Patent No. 5942430  
; GENERAL INFORMATION:  
; APPLICANT: ROBERTSON, Daniel E.  
; APPLICANT: MURPHY, Dennis  
; APPLICANT: REID, John  
; APPLICANT: MAFFIA, Anthony  
; APPLICANT: LINK, Steven  
; APPLICANT: SWANSON, Ronald V.  
; APPLICANT: WARREN, Patrick V.

APPLICANT: KOSMOTKA, Anna  
TITLE OF INVENTION: ESTERASES  
NUMBER OF SEQUENCES: 42  
CORRESPONDENCE ADDRESS:  
ADDRESSEE: FISH & RICHARDSON P.C.  
STREET: 4225 EXECUTIVE SQUARE, STE 1400  
CITY: LA JOLLA  
STATE: CALIFORNIA  
COUNTRY: USA

ZIP: 92037  
COMPUTER READABLE FORM:  
MEDIUM TYPE: 3.5 INCH DISKETTE  
COMPUTER: IBM PS/2  
OPERATING SYSTEM: MS-DOS  
SOFTWARE: WORD PERFECT 6.0  
CURRENT APPLICATION DATA:  
APPLICATION NUMBER: US/08/602,359A  
FILING DATE: February 16, 1996  
CLASSIFICATION: 435  
PRIOR APPLICATION DATA:  
APPLICATION NUMBER:  
FILING DATE:  
ATTORNEY/AGENT INFORMATION:  
NAME: HAILE, LISA A.  
REGISTRATION NUMBER: 38,347  
REFERENCE/DOCKET NUMBER: 09010/010001  
TELECOMMUNICATION INFORMATION:  
TELEPHONE: 619-678-5070  
TELEFAX: 619-678-5099  
INFORMATION FOR SEQ ID NO: 31:  
SEQUENCE CHARACTERISTICS:  
LENGTH: 936 NUCLEOTIDES  
TYPE: NUCLEIC ACID  
STRANDEDNESS: SINGLE  
TOPOLOGY: LINEAR  
MOLECULE TYPE: GENOMIC DNA  
US-08-602-359A-31

Query Match 22.0%; Score 28.4; DB 2; Length 936;  
Best Local Similarity 56.4%; Pred. No. 2.8;  
Matches 53; Conservative 0; Mismatches 41; Indels 0; Gaps 0;  
  
QY 2 GACCTTCTGCTCGTAGCGATTACTTCGAGCAATTAAGCAAGACCGGACCGAGAT 61  
DB 718 GACCTTGAACCTACTCTCTGCGCTGATATACCGCGGATACGACCGGCTGAGAT 777  
  
QY 62 GGTGGGGTCTTTTGTGTGTGCTGTGAGTG 95  
DB 778 GAAGGAGAAGTTTTCGGGCGAGATGCTGAGAAGAG 811

RESULT 10  
US-09-270-767-13705/c  
Sequence 13705, Application US/09270767  
Patent No. 6703491  
GENERAL INFORMATION:  
APPLICANT: Homburger et al.  
TITLE OF INVENTION: Nucleic acids and proteins of *Drosophila melanogaster*  
FILE REFERENCE: File Reference: 7326-094  
CURRENT APPLICATION NUMBER: US/09/270,767  
CURRENT FILING DATE: 1999-03-17  
NUMBER OF SEQ ID NOS: 62517  
SOFTWARE: PatentIn Ver. 2.0  
SEQ ID NO 13705  
LENGTH: 1160  
TYPE: DNA  
ORGANISM: *Drosophila melanogaster*  
US-09-270-767-13705

Query Match 21.2%; Score 27.4; DB 3; Length 1160;  
Best Local Similarity 54.5%; Pred. No. 7.2;  
Matches 55; Conservative 0; Mismatches 46; Indels 0; Gaps 0;  
  
QY 4 CTTTCTGCTCGTAGCGATTACTTCGAGCAATTAAGCAAGACCGGACCGAGATGG 63  
DB 656 CCATTTTCGAGTCGTCGCTAATTCATCCACCTTGTTGTTAATACACCTCGAACTTGATGC 597  
  
QY 64 TCGGGCTCTTTTGTGTGCTGTGAGCGTGTGTCCTCAAC 104  
DB 596 GCTTGTTCACCTTGTAGGCTGCAGCACTGTGTCTTGAAC 556

RESULT 11  
US-10-144-929-56

Sequence 56, Application US/10144929  
Patent No. 6881823  
GENERAL INFORMATION:  
APPLICANT: Ruben et al.  
TITLE OF INVENTION: 70 Human Secreted Proteins  
FILE REFERENCE: P2014P1  
CURRENT APPLICATION NUMBER: US/10/144,929  
CURRENT FILING DATE: 2002-05-15  
PRIOR APPLICATION NUMBER: US/09/251,329  
PRIOR FILING DATE: 1999-02-17  
PRIOR APPLICATION NUMBER: PCT/US98/17044  
PRIOR FILING DATE: 1998-08-18  
NUMBER OF SEQ ID NOS: 257  
SOFTWARE: PatentIn Ver. 2.0  
SEQ ID NO 56  
LENGTH: 1951  
TYPE: DNA  
ORGANISM: Homo sapiens  
FEATURE:  
NAME/KEY: SITE  
LOCATION: (28)  
OTHER INFORMATION: n equals a,t,g, or c  
FEATURE:  
NAME/KEY: SITE  
LOCATION: (1636)  
OTHER INFORMATION: n equals a,t,g, or c  
FEATURE:  
NAME/KEY: SITE  
LOCATION: (1947)  
OTHER INFORMATION: n equals a,t,g, or c  
FEATURE:  
NAME/KEY: SITE  
LOCATION: (1951)  
OTHER INFORMATION: n equals a,t,g, or c  
US-10-144-929-56

Query Match 21.2%; Score 27.4; DB 3; Length 1951;  
Best Local Similarity 54.5%; Pred. No. 8.9;  
Matches 55; Conservative 0; Mismatches 46; Indels 0; Gaps 0;  
  
QY 24 CTTGAGCAATTAAGCAAGACCGGACCGGATGTCGGGCTCTTTTGTGTTGG 83  
DB 718 CTGCTGGGATCTTCGTTGGATTAATCCCTCACCTCTCGGCGATGTGTTTCTTGTGG 777  
  
QY 84 TGCTGTGAGTGTGTCACCGTATTATTTCGGACTAGTT 124  
DB 778 GGTGTAACTGCTGCGCCACTTCAATGCTACTCTGT 818

RESULT 12  
US-09-949-016-125432/c  
Sequence 125432, Application US/09949016  
Patent No. 6812339  
GENERAL INFORMATION:  
APPLICANT: VENTER, J. Craig et al.  
TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED  
WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF  
FILE REFERENCE: CL001307  
CURRENT APPLICATION NUMBER: US/09/949,016  
CURRENT FILING DATE: 2000-04-14  
PRIOR APPLICATION NUMBER: 60/241,755  
PRIOR FILING DATE: 2000-10-20

;; PRIOR APPLICATION NUMBER: 60/237,768  
;; PRIOR FILING DATE: 2000-10-03  
;; PRIOR APPLICATION NUMBER: 60/231,498  
;; PRIOR FILING DATE: 2000-09-08  
;; NUMBER OF SEQ ID NOS: 207012  
;; SOFTWARE: FastSeq for Windows Version 4.0  
;; SEQ ID NO 125432  
;; LENGTH: 601  
;; TYPE: DNA  
;; ORGANISM: Human  
US-09-949-016-125432

Query Match 20.6%; Score 26.6; DB 3; Length 601;  
Best Local Similarity 66.7%; Pred. No. 11;  
Matches 38; Conservative 0; Mismatches 19; Indels 0; Gaps 0;  
QY 44 AAAGACCCGACGAGATGTCGGGGCTTTTGTGTGCTGTGACGTGTGTC 100  
DB 407 AAAGCCTCTGATTGAGATGAGGAAGGTCTTCTACCTCTGAGCTGTAAGGTCTGTGC 351

RESULT 13  
US-09-949-016-15300  
;; Sequence 15300, Application US/09949016  
;; Patent No. 6812339  
;; GENERAL INFORMATION:  
;; APPLICANT: VENTER, J. Craig et al.  
;; TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED  
;; TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF  
;; FILE REFERENCE: CL001307  
;; CURRENT APPLICATION NUMBER: US/09/949,016  
;; CURRENT FILING DATE: 2000-04-14  
;; PRIOR APPLICATION NUMBER: 60/241,755  
;; PRIOR FILING DATE: 2000-10-20  
;; PRIOR APPLICATION NUMBER: 60/237,768  
;; PRIOR FILING DATE: 2000-10-03  
;; PRIOR APPLICATION NUMBER: 60/231,498  
;; PRIOR FILING DATE: 2000-09-08  
;; NUMBER OF SEQ ID NOS: 207012  
;; SOFTWARE: FastSeq for Windows Version 4.0  
;; SEQ ID NO 15300  
;; LENGTH: 139552  
;; TYPE: DNA  
;; ORGANISM: Human  
US-09-949-016-15300

Query Match 20.6%; Score 26.6; DB 3; Length 139552;  
Best Local Similarity 66.7%; Pred. No. 99;  
Matches 38; Conservative 0; Mismatches 19; Indels 0; Gaps 0;  
QY 44 AAAGACCCGACGAGATGTCGGGGCTTTTGTGTGCTGTGACGTGTGTC 100  
DB 80719 AAAGCCTCTGATTGAGATGAGGAAGGTCTTCTACCTCTGAGCTGTAAGGTCTGTGC 80775

RESULT 14  
US-09-312-762A-3/c  
;; Sequence 3, Application US/09312762A  
;; Patent No. 6552177  
;; GENERAL INFORMATION:  
;; APPLICANT: MIA HOROWITZ ET AL.  
;; TITLE OF INVENTION: EH DOMAIN CONTAINING GENES AND PROTEINS  
;; NUMBER OF SEQUENCES: 27  
;; CORRESPONDENCE ADDRESS:  
;; ADDRESSEE: Mark M. Friedman c/o Anthony Castorina  
;; STREET: 2001 Jefferson Davis Highway, Suite 207  
;; CITY: Arlington  
;; STATE: Virginia  
;; COUNTRY: United States of America  
;; ZIP: 22202  
;; COMPUTER READABLE FORM:  
;; MEDIUM TYPE: 1.44 megabyte, 3.5" microdisk  
;; COMPUTER: Twinhead\* Slimnote-890TX

;; OPERATING SYSTEM: MS DOS version 6.2,  
;; OPERATING SYSTEM: Windows version 3.11  
;; SOFTWARE: Word for Windows version 2.0 converted to  
;; SOFTWARE: an ASCII file  
;; CURRENT APPLICATION DATA:  
;; APPLICATION NUMBER: US/09/312,762A  
;; FILING DATE:  
;; CLASSIFICATION:  
;; PRIOR APPLICATION DATA:  
;; APPLICATION NUMBER: 09/026,898  
;; FILING DATE: 20 FEB 1998  
;; ATTORNEY/AGENT INFORMATION:  
;; NAME: Friedmam, Mark M.  
;; REGISTRATION NUMBER: 33,883  
;; REFERENCE/DOCKET NUMBER: 916/10  
;; TELECOMMUNICATION INFORMATION:  
;; TELEPHONE: 972-3-5625553  
;; TELEFAX: 972-3-5625554  
;; TELEX:  
;; INFORMATION FOR SEQ ID NO: 3:  
;; SEQUENCE CHARACTERISTICS:  
;; LENGTH: 14707  
;; TYPE: nucleic acid  
;; STRANDEDNESS: double  
;; TOPOLOGY: linear  
US-09-312-762A-3

Query Match 20.5%; Score 26.4; DB 3; Length 14707;  
Best Local Similarity 54.0%; Pred. No. 47;  
Matches 54; Conservative 0; Mismatches 46; Indels 0; Gaps 0;  
QY 7 TCTGCTCGTAGCGATTACTTCGAGCAATTACTGACGCAAAAGACCCGACGAGTGTGCG 66  
DB 5137 TCTGTCTCTGCTGATTTTGGGGCTTGTGATGACTTCTGAGAACTGCTCTGAGATGTCCA 5078  
QY 67 GGGTCTTTTGTGTGCTGTGACGTGTGTCTCCACCG 106  
DB 5077 GCTTGTGGCGTCGAAACAAAGAGATCGGTCCACAG 5038

RESULT 15  
US-09-949-016-143314/c  
;; Sequence 143314, Application US/09949016  
;; Patent No. 6812339  
;; GENERAL INFORMATION:  
;; APPLICANT: VENTER, J. Craig et al.  
;; TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED  
;; TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF  
;; FILE REFERENCE: CL001307  
;; CURRENT APPLICATION NUMBER: US/09/949,016  
;; CURRENT FILING DATE: 2000-04-14  
;; PRIOR APPLICATION NUMBER: 60/241,755  
;; PRIOR FILING DATE: 2000-10-20  
;; PRIOR APPLICATION NUMBER: 60/237,768  
;; PRIOR FILING DATE: 2000-10-03  
;; PRIOR APPLICATION NUMBER: 60/231,498  
;; PRIOR FILING DATE: 2000-09-08  
;; NUMBER OF SEQ ID NOS: 207012  
;; SOFTWARE: FastSeq for Windows Version 4.0  
;; SEQ ID NO 143314  
;; LENGTH: 601  
;; TYPE: DNA  
;; ORGANISM: Human  
US-09-949-016-143314

Query Match 20.3%; Score 26.2; DB 3; Length 601;  
Best Local Similarity 58.2%; Pred. No. 15;  
Matches 46; Conservative 0; Mismatches 33; Indels 0; Gaps 0;  
QY 29 AGCATTTACTGACGACAAAGACCCCGACGAGATGTCGGGGCTTTTGTGTGTTGTTGTCG 88  
DB 291 AGCTTCTCTGACACCCACCCACCCAGTCTTGTGTGTTGTTGTTGTTGTTGTTGTTGTTG 232

Qy 89 TGACGTGTGTCTCAACCGT 107  
||| ||| ||| ||| ||| |||  
Db 231 TGGCTTTTGTTCACCTGT 213

Search completed: March 7, 2006, 14:11:48  
Job time : 93.4206 secs



GenCore version 5.1.7  
Copyright (c) 1993 - 2006 Bioceleration Ltd.

OM nucleic - nucleic search, using sw model

Run on: March 7, 2006, 12:18:50 ; Search time 1192.75 Seconds  
(without alignments)  
237.125 Million cell updates/sec

Title: US-10-782-899-1\_COPY\_472\_600

Perfect score: 129  
Sequence: 1 tgacctctgctgcagcga.....tattccggactagtcagcg 129

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 7218535 seqs, 1096242582 residues

Total number of hits satisfying chosen parameters: 14437070

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published Applications\_NA\_New.\*  
1: /cgn2\_6/ptodata/1/pubpna/US08\_NEW\_PUB.seq.\*  
2: /cgn2\_6/ptodata/1/pubpna/US06\_NEW\_PUB.seq.\*  
3: /cgn2\_6/ptodata/1/pubpna/US07\_NEW\_PUB.seq.\*  
4: /cgn2\_6/ptodata/1/pubpna/US09\_NEW\_PUB.seq.\*  
5: /cgn2\_6/ptodata/1/pubpna/US09\_NEW\_PUB.seq.\*  
6: /cgn2\_6/ptodata/1/pubpna/US09\_NEW\_PUB.seq.\*  
7: /cgn2\_6/ptodata/1/pubpna/US10\_NEW\_PUB.seq.\*  
8: /cgn2\_6/ptodata/1/pubpna/US10\_NEW\_PUB.seq.\*  
9: /cgn2\_6/ptodata/1/pubpna/US11\_NEW\_PUB.seq.\*  
10: /cgn2\_6/ptodata/1/pubpna/US11\_NEW\_PUB.seq.\*  
11: /cgn2\_6/ptodata/1/pubpna/US11\_NEW\_PUB.seq.\*  
12: /cgn2\_6/ptodata/1/pubpna/US11\_NEW\_PUB.seq.\*  
13: /cgn2\_6/ptodata/1/pubpna/US60\_NEW\_PUB.seq.\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	29	22.5	185750	US-11-114-798-56	Sequence 56, Appl
2	29	22.5	1080000	US-10-928-446A-1	Sequence 1, Appl1
3	29	22.5	1080000	US-10-928-446A-181	Sequence 181, App
4	29	22.5	1080000	US-10-928-446A-183	Sequence 183, App
5	29	22.5	1080000	US-10-928-446A-185	Sequence 185, App
6	29	22.5	1080000	US-10-928-446A-187	Sequence 187, App
7	29	22.5	1080000	US-10-928-446A-189	Sequence 189, App
8	29	22.5	1080000	US-10-928-446A-191	Sequence 191, App
9	29	22.5	1080000	US-10-928-446A-193	Sequence 193, App
10	29	22.5	1080000	US-10-928-446A-195	Sequence 195, App
11	29	22.5	1080000	US-10-928-446A-197	Sequence 197, App
12	29	22.5	1080000	US-10-928-446A-199	Sequence 199, App
13	29	22.5	1080000	US-10-928-446A-201	Sequence 201, App
14	28.8	22.3	79134	US-11-124-368A-2924	Sequence 2924, Ap
15	27.6	21.4	2150	US-10-955-054A-98	Sequence 98, Appl
16	27.6	21.4	2150	US-11-072-175-35	Sequence 35, Appl
17	27.4	21.2	398	US-11-000-688-1241	Sequence 1241, Ap
18	27.4	21.2	1356	US-10-973-115B-43	Sequence 43, Appl
19	27.4	21.2	1356	US-10-131-826A-43	Sequence 43, Appl
20	26.4	20.5	523	US-09-925-065A-108531	Sequence 108531,

21	26.4	20.5	1223	6	US-09-925-065A-694721	Sequence 694721,
22	26.4	20.5	1223	6	US-09-925-065A-694722	Sequence 694722,
23	26.4	20.5	2499	8	US-10-750-185-28315	Sequence 28315, A
24	26.4	20.5	2499	8	US-10-750-623-28315	Sequence 28315, A
25	26.2	20.3	495	6	US-09-925-065A-684535	Sequence 684535,
26	26.2	20.3	495	6	US-09-925-065A-684536	Sequence 684536,
27	26.2	20.3	578	12	US-11-128-061-1368	Sequence 1368, Ap
28	26.2	20.3	578	12	US-11-128-061-5010	Sequence 5010, Ap
29	26.2	20.3	578	12	US-11-128-049-1368	Sequence 1368, Ap
30	26.2	20.3	578	12	US-11-128-049-5010	Sequence 5010, Ap
31	26.2	20.3	121736	9	US-11-114-798-49	Sequence 49, Appl
32	26.2	20.3	318488	9	US-11-114-798-58	Sequence 58, Appl
33	26	20.2	565	6	US-09-925-065A-663286	Sequence 663286,
34	26	20.2	565	6	US-09-925-065A-663287	Sequence 663287,
35	26	20.2	600	12	US-11-136-527-7332	Sequence 7332, Ap
36	26	20.2	1025	12	US-11-136-527-3436	Sequence 3436, Ap
37	26	20.2	1426	6	US-09-925-065A-84896	Sequence 84896, A
38	26	20.2	1617	6	US-09-925-065A-81196	Sequence 81196, A
39	26	20.2	33931	12	US-11-124-367A-5006	Sequence 5006, Ap
40	25.8	20.0	1808	6	US-09-925-065A-679127	Sequence 679127,
41	25.6	19.8	463	6	US-09-925-065A-476394	Sequence 476394,
42	25.6	19.8	560	6	US-09-925-065A-251626	Sequence 251626,
43	25.4	19.7	443	6	US-09-925-065A-239621	Sequence 239621,
44	25.4	19.7	567	6	US-09-925-065A-770926	Sequence 770926,
45	25.4	19.7	2238	8	US-10-750-185-40094	Sequence 40094, A

ALIGNMENTS

RESULT 1

US-11-114-798-56  
; Sequence 56, Application US/11114798  
; Publication No. US20060035246A1  
; GENERAL INFORMATION:  
; APPLICANT: WU, RINA  
; APPLICANT: MARQUEZ, ABBEY  
; TITLE OF INVENTION: CHROMOGENIC IN SITU HYBRIDIZATION METHODS, KITS, AND  
; FILE REFERENCE: 0618.011.0004  
; CURRENT APPLICATION NUMBER: US/11/114,798  
; CURRENT FILING DATE: 2005-04-26  
; PRIOR FILING DATE: 2002-06-17  
; PRIOR APPLICATION NUMBER: 10/173,525  
; PRIOR FILING DATE: 2001-09-14  
; PRIOR APPLICATION NUMBER: 09/952,851  
; NUMBER OF SEQ ID NOS: 58  
; SOFTWARE: PatentIn Ver. 3.3  
; SEQ ID NO 56  
; LENGTH: 185750  
; TYPE: DNA  
; ORGANISM: Artificial Sequence  
; FEATURE:  
; OTHER INFORMATION: Description of Artificial Sequence: Synthetic  
; OTHER INFORMATION: nucleotide sequence  
US-11-114-798-56

Query Match	22.5%;	Score 29;	DB 9;	Length 185750;
Best Local Similarity	55.4%;	Pred. No. 7.9;		
Matches	56;	Conservative	0;	Mismatches 45; Indels 0; Gaps 0;
QY	16	AGCGATTACTTTCAGCATTACTGACGACAAAGACCCGACGAGATGGTCGGGTCCTTTT	75	
Db	70115	AGCAGTTCACTTGAGACTTCTCAGCGAAGAGTGGGATACAGTTGGTGTAGTCTTAT	70174	
QY	76	TGTTGGTGGTGTGACGTTGTGTCTCAACGGTATTATTCG	116	
Db	70175	ATTGTAGCTGCTCATGTGTCTCTGATGATAGATGCAG	70215	

RESULT 2

US-10-928-446A-1  
; Sequence 1, Application US/10928446A

```
; Publication No. US20050277123A1
; GENERAL INFORMATION:
; APPLICANT: UNIVERSITY OF UTAH RESEARCH FOUNDATION
; TITLE OF INVENTION: VARIANTS OF NEDD4L ASSOCIATED WITH HYPERTENSION AND
; TITLE OF INVENTION: VIRAL BUDDING
; FILE REFERENCE: 0274-5785.1US
; CURRENT APPLICATION NUMBER: US/10/928,446A
; CURRENT FILING DATE: 2004-08-26
; PRIOR APPLICATION NUMBER: 60/359,741
; PRIOR FILING DATE: 2002-02-26
; NUMBER OF SEQ ID NOS: 202
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 1
; LENGTH: 1080000
; TYPE: DNA
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: allele
; LOCATION: (825234)
; OTHER INFORMATION: the 'n' at position 825234 may be 'c' or 't'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (825270)
; OTHER INFORMATION: the 'n' at position 825270 may be 'c' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (825401)
; OTHER INFORMATION: the 'n' at position 825401 may be 'c' or 'a'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (825428)
; OTHER INFORMATION: the 'n' at position 825428 may be 'g' or 'a'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (825473)
; OTHER INFORMATION: the 'n' at position 825473 may be 'g' or 'a'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (825765)
; OTHER INFORMATION: a "c" may be deleted at this position
; FEATURE:
; NAME/KEY: allele
; LOCATION: (825828)
; OTHER INFORMATION: the 'n' at position 825828 may be 'c' or 't'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (826041)
; OTHER INFORMATION: the 'n' at position 826041 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (826546)
; OTHER INFORMATION: the 'n' at position 826546 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (826654)
; OTHER INFORMATION: the 'n' at position 826654 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (826826)
; OTHER INFORMATION: the 'n' at position 826826 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (826863)
; OTHER INFORMATION: the 'n' at position 826863 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (827008)
; OTHER INFORMATION: the 'n' at position 827008 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (843055)
; OTHER INFORMATION: the 'n' at position 843055 may be 'g' or 'a'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (843118)
; OTHER INFORMATION: the 'n' at position 843118 may be 't' or 'c'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (871027)
; OTHER INFORMATION: the 'n' at position 871027 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (871140)
; OTHER INFORMATION: the 'n' at position 871140 may be 'c' or 't'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (871168)
; OTHER INFORMATION: the 'n' at position 871168 may be 'c' or 'a'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (872678)
; OTHER INFORMATION: the 'n' at position 872678 may be 'c' or 't'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (872742)
; OTHER INFORMATION: the 'n' at position 872742 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (925859)
; OTHER INFORMATION: the 'n' at position 925859 may be 'c' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (993220)
; OTHER INFORMATION: the 'n' at position 993220 may be 'c' or 't'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (993254)
; OTHER INFORMATION: the 'n' at position 993254 may be 'g' or 'a'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1006462)
; OTHER INFORMATION: the 'n' at position 1006462 may be 'c' or 't'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1007820)..(1007823)
; OTHER INFORMATION: "ttct" may be deleted at this position
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1018038)
; OTHER INFORMATION: the 'n' at position 1018038 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1018704)
; OTHER INFORMATION: the 'n' at position 1018704 may be 'c' or 't'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1018718)..(1018720)
; OTHER INFORMATION: "gtt" may be deleted at this position
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1026786)
; OTHER INFORMATION: the 'n' at position 1026786 may be 'c' or 'a'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1047134)
; OTHER INFORMATION: the 'n' at position 1047134 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1047159)
; OTHER INFORMATION: the 'n' at position 1047159 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1047378)
; OTHER INFORMATION: the 'n' at position 1047378 may be 'c' or 't'
; FEATURE:
; NAME/KEY: allele
```

```
; LOCATION: (1047739)
; OTHER INFORMATION: the 'n' at position 1047739 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1050133)..(1050137)
; OTHER INFORMATION: "ctaaa" may be deleted at this position
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1050539)
; OTHER INFORMATION: the 'n' at position 1050539 may be 'c' or 't'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1062808)
; OTHER INFORMATION: the 'n' at position 1062808 may be 'c' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1066392)
; OTHER INFORMATION: the 'n' at position 1066392 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: allele
; LOCATION: (1073711)
; OTHER INFORMATION: the 'n' at position 1073711 may be 'c' or 't'
;
US-10-928-446A-1
Query Match 22.5%; Score 29; DB 8; Length 1080000;
Best Local Similarity 55.4%; Pred. No. 12;
Matches 56; Conservative 0; Mismatches 45; Indels 0; Gaps 0;

Qy 16 AGCGATTACTCGAGCATTTACTGACGACAAAGACCCGACGATGCGGGTCTTTT 75
Db 771426 AGCAGTTCACTTGAGACTTTTCTACCGAAAGAGTGCATACAGTTGGTGTAGTCTTAT 771485

Qy 76 TGTGTGGTGCTGTGAGCTGTGTCCAAACCGTATTATTCGG 116
Db 771486 ATTGTACTGCTCATGTGTGTCATGATAGATGCAG 771526

RESULT 3
US-10-928-446A-181
; Sequence 181, Application US/10928446A
; Publication No. US2005027123A1
; GENERAL INFORMATION:
; APPLICANT: UNIVERSITY OF UTAH RESEARCH FOUNDATION
; TITLE OF INVENTION: VARIANTS OF NEDD4L ASSOCIATED WITH HYPERTENSION AND
; TITLE OF INVENTION: VIRAL BUDDING
; FILE REFERENCE: 0274-5785.1US
; CURRENT APPLICATION NUMBER: US/10/928,446A
; CURRENT FILING DATE: 2004-08-26
; PRIOR APPLICATION NUMBER: 60/359,741
; PRIOR FILING DATE: 2002-02-26
; NUMBER OF SEQ ID NOS: 202
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 181
; LENGTH: 1080000
; TYPE: DNA
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (826985)..(827008)
; FEATURE:
; OTHER INFORMATION: full exon 1 range is 826667-827008
; FEATURE:
; NAME/KEY: allele
; LOCATION: (827008)..(827008)
; OTHER INFORMATION: the 'n' at position 827008 may be 'a' or 'g'
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (843242)..(843315)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (922549)..(922630)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (926021)..(926059)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (929123)..(929176)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (993104)..(993154)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (999547)..(999608)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1000354)..(1000456)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1002118)..(1002284)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1006117)..(1006249)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1007860)..(1008036)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1010940)..(1011014)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1018160)..(1018291)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1018800)..(1018919)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1020028)..(1020225)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1026659)..(1026736)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1028113)..(1028167)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1034316)..(1034374)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1041390)..(1041455)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1043121)..(1043350)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1044868)..(1044989)
; OTHER INFORMATION: exon
; FEATURE:
```

```
; NAME/KEY: CDS
; LOCATION: (1047519)..(1047589)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1050296)..(1050391)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1060368)..(1060441)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1062648)..(1062708)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1064561)..(1064620)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1066207)..(1066314)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1067768)..(1067864)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1068609)..(1068681)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1073289)..(1073388)
; FEATURE:
; OTHER INFORMATION: full exon 30 range is 1073289-1075279
US-10-928-446A-181

Query Match      22.5%; Score 29; DB 8; Length 1080000;
Best Local Similarity 55.4%; Pred. No. 12;
Matches 56; Conservative 0; Mismatches 45; Indels 0; Gaps 0;

QY      16  AGCGATTACTTCGAGCATTACTGACGACAAAGACCCGACCGAGATGGTCGGGTCTTTT 75
Db      771426  AGCAGTTCATTGAGACTTTCTCAGGAAGAGTGCATGTTGGTAGTCTTAT 771485

QY      76  TGTGTGGTGTGTGACGTGTGTGCCAACCGTATATTCCG 116
Db      771486  ATTGTGCTACGTGCTCATGTGTGTCTGATGATGAATGCAG 771526

RESULT 4
US-10-928-446A-183
; Sequence 183, Application US/10928446A
; Publication No. US20050277123A1
; GENERAL INFORMATION:
; APPLICANT: UNIVERSITY OF UTAH RESEARCH FOUNDATION
; TITLE OF INVENTION: VARIANTS OF NEDD4L ASSOCIATED WITH HYPERTENSION AND
; TITLE OF INVENTION: VIRAL BUDDING
; FILE REFERENCE: 0274-5785.1US
; CURRENT APPLICATION NUMBER: US/10/928,446A
; CURRENT FILING DATE: 2004-08-26
; PRIOR APPLICATION NUMBER: 60/359,741
; PRIOR FILING DATE: 2002-02-26
; NUMBER OF SEQ ID NOS: 202
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 183
; LENGTH: 1080000
; TYPE: DNA
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (826985)..(827017)
; FEATURE:
; OTHER INFORMATION: full exon 1 range is 826667-827008
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (843242)..(843315)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (922549)..(922630)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (926021)..(926059)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (929123)..(929176)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (993104)..(993154)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (999547)..(999608)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1000354)..(1000456)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1002118)..(1002284)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1006117)..(1006249)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1007860)..(1008036)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1010940)..(1011014)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1018160)..(1018291)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1018800)..(1018919)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1020028)..(1020225)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1026659)..(1026736)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1028113)..(1028167)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1034316)..(1034374)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1041390)..(1041455)
```

```
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1043121)..(1043350)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1044868)..(1044989)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1047519)..(1047589)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1050296)..(1050391)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1060368)..(1060441)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1062648)..(1062708)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1064561)..(1064620)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1066207)..(1066314)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1067768)..(1067864)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1068609)..(1068681)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1073289)..(1073388)
OTHER INFORMATION: full exon 30 range is 1073289-1075279
US-10-928-446A-183
```

```
Query Match 22.5%; Score 29; DB 8; Length 1080000;
Best Local Similarity 55.4%; Pred. No. 12;
Matches 56; Conservative 0; Mismatches 45; Indels 0; Gaps 0;

Qy 16 AGCGATTACTCGAGCTTACTGACGACAAGACCCGACGAGATGTCGGGCTCTTT 75
Db 771426 AGCAGTTCACTTGAGACTTTCTCACGGAAGAAGTGCATGACATGTCGTAT 771485
Qy 76 TGTGTGCTGTGAGTGTTGCCAACCGTATTATTCG 116
Db 771486 ATTGCTACGTGCTCATGTGTCTGATGATGATGACG 771526
```

```
RESULT 5
US-10-928-446A-185
Sequence 185 Application US/10928446A
Publication No. US2005027123A1
GENERAL INFORMATION:
APPLICANT: UNIVERSITY OF UTAH RESEARCH FOUNDATION
TITLE OF INVENTION: VARIANTS OF NEDD4L ASSOCIATED WITH HYPERTENSION AND
TITLE OF INVENTION: VIRAL BUDDING
FILE REFERENCE: 0274-5785.1US
CURRENT APPLICATION NUMBER: US/10/928,446A
CURRENT FILING DATE: 2004-08-26
PRIOR APPLICATION NUMBER: 60/359,741
```

```
PRIOR FILING DATE: 2002-02-26
NUMBER OF SEQ ID NOS: 202
SOFTWARE: PatentIn Ver. 2.1
SEQ ID NO 185
LENGTH: 1080000
TYPE: DNA
ORGANISM: Homo sapiens
FEATURE:
NAME/KEY: CDS
LOCATION: (722487)..(722534)
FEATURE:
OTHER INFORMATION: full length exon 1 range is 722213-722534
FEATURE:
NAME/KEY: CDS
LOCATION: (843242)..(843315)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (922549)..(922630)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (926021)..(926059)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (929123)..(929176)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (993104)..(993154)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (999547)..(999608)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1000354)..(1000456)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1002118)..(1002284)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1006117)..(1006249)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1007860)..(1008036)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1010940)..(1011014)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1018160)..(1018291)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1018800)..(1018919)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1020028)..(1020225)
OTHER INFORMATION: exon
FEATURE:
NAME/KEY: CDS
LOCATION: (1026659)..(1026736)
OTHER INFORMATION: exon
FEATURE:
```



```
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1044868)..(1044989)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1047519)..(1047589)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1050296)..(1050391)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1060368)..(1060441)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1062648)..(1062708)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1064561)..(1064620)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1066207)..(1066314)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1067768)..(1067864)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1068609)..(1068681)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1073289)..(1073388)
/ OTHER INFORMATION: full length exon 30 range is 1073289-1075279
/ US-10-928-446A-187
Query Match 22.5%; Score 29; DB 8; Length 1080000;
Best Local Similarity 55.4%; Pred. No. 12;
Matches 56; Conservative 0; Mismatches 45; Indels 0; Gaps 0;
QY 16 AGCGATTACTTCGAGCATTACTGACGACAAAGACCCGACGAGATGGTCGGGCTCTTTT 75
Db 771426 AGCAGTTCACCTCGAGACTTCTCACGGAAGAAGTGCATACAGTTGGTGTAGTCTTAT 771485
QY 76 TGTGTGGTCTGTGAGGTGTTGTCACCGTATTATTCCG 116
Db 771486 ATTGTGCTACGTGCTGTGTTGTTCTGATGATAGAATGCAG 771526
RESULT 7
US-10-928-446A-189
; Sequence 189, Application US/10928446A
; Publication No. US2005027123A1
; GENERAL INFORMATION:
; APPLICANT: UNIVERSITY OF UTAH RESEARCH FOUNDATION
; TITLE OF INVENTION: VARIANTS OF NEDD4L ASSOCIATED WITH HYPERTENSION AND
; TITLE OF INVENTION: VIRAL BUDDING
; FILE REFERENCE: 0274-5785.IUS
; CURRENT APPLICATION NUMBER: US/10/928,446A
; CURRENT FILING DATE: 2004-08-26
; PRIOR APPLICATION NUMBER: 60/359,741
; PRIOR FILING DATE: 2002-02-26
; NUMBER OF SEQ ID NOS: 202
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 189
/ LENGTH: 1080000
/ TYPE: DNA
/ ORGANISM: Homo sapiens
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (723508)..(723516)
/ FEATURE:
/ OTHER INFORMATION: full length exon 1 range is 723483-723516
/ NAME/KEY: CDS
/ LOCATION: (1000354)..(1000456)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1002118)..(1002284)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1006117)..(1006249)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1007860)..(1008036)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1010940)..(1011014)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1018160)..(1018291)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1018800)..(1018919)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1020028)..(1020225)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1026659)..(1026736)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1028113)..(1028167)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1034316)..(1034374)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1041390)..(1041455)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1043121)..(1043350)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1044868)..(1044989)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1047519)..(1047589)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1050296)..(1050391)
/ OTHER INFORMATION: exon
/ FEATURE:
```

```
/ NAME/KEY: CDS
/ LOCATION: (1060368)..(1060441)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1062648)..(1062708)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1064561)..(1064620)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1066207)..(1066314)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1067768)..(1067864)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1068609)..(1068681)
/ OTHER INFORMATION: exon
/ FEATURE:
/ NAME/KEY: CDS
/ LOCATION: (1073289)..(1073388)
/ OTHER INFORMATION: full length exon 30 range is 1073289-1075279
/ OTHER INFORMATION: full length exon 30 range is 1073289-1075279
US-10-928-446A-189

Query Match 22.5% Score 29; DB 8; Length 1080000;
Best Local Similarity 55.4%; Pred. No. 12; Mismatches 45; Indels 0; Gaps 0;
Matches 56; Conservative 0;

QY 16 AGCGATTACTTCGAGCATTAAGCGACAAAGACCCGACGAGATGGTCGGGTCTTTT 75
Db 771426 AGCAGTTCACTTCGAGACTTCTCAGCGAAGAGTGCGATACAGTTGGTGTAGTCTTAT 771485
QY 76 TGTGTGGTCTGTGAGCTGTGTCACCAACCGTATTATTCG 116
Db 771486 ATTGTGCTACGTGCTGTGTTGCTGTGATGATGAATGCAG 771526

RESULT 8
US-10-928-446A-191
; Sequence 191, Application US/10928446A
; Publication No. US2005027123A1
; GENERAL INFORMATION:
; APPLICANT: UNIVERSITY OF UTAH RESEARCH FOUNDATION
; TITLE OF INVENTION: VARIANTS OF NEDD4L ASSOCIATED WITH HYPERTENSION AND
; TITLE OF INVENTION: VIRAL BUDDING
; FILE REFERENCE: 0274-5785.1US
; CURRENT APPLICATION NUMBER: US/10/928,446A
; CURRENT FILING DATE: 2004-08-26
; PRIOR APPLICATION NUMBER: 60/359,741
; PRIOR FILING DATE: 2002-02-26
; NUMBER OF SEQ ID NOS: 202
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 191
; LENGTH: 1080000
; TYPE: DNA
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (999562)..(999608)
; FEATURE:
; OTHER INFORMATION: full length exon 7 range is 999547-999608
; NAME/KEY: CDS
; LOCATION: (1000354)..(1000456)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1002118)..(1002284)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1006117)..(1006249)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1007860)..(1008036)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1010940)..(1011014)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1018160)..(1018291)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1018800)..(1018919)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1020028)..(1020225)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1026659)..(1026736)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1028113)..(1028167)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1034316)..(1034374)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1041390)..(1041455)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1043121)..(1043350)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1044868)..(1044989)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1047519)..(1047589)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1050296)..(1050391)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1060368)..(1060441)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1062648)..(1062708)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1064561)..(1064620)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1066207)..(1066314)
```



```
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1067768)..(1067864)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1068609)..(1068681)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1073289)..(1073388)
; FEATURE:
; OTHER INFORMATION: full length exon 30 range is 1073289-1075279
US-10-928-446A-191

Query Match          22.5%  Score 29;  DB 8;  Length 1080000;
Best Local Similarity 55.4%  Pred.No. 12;
Matches 56;  Conservative 0;  Mismatches 45;  Indels 0;  Gaps 0;

QY 16 AGCGATTACTTCGAGCATTACTGACGACAAAGACCCCGACGAGATGGTCGGGTCCTTTT 75
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 771426 AGCAGTTCACTTGAGACTTCTCACGGAAGAGTGCATACAGTTGGTGTAGTCTTAT 771485

QY 76 TGTGTGGTGTGTGACGTGTGTGTCACACCGTATTATTCCG 116
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 771486 ATTTGCTACGTGCTCATGTGTGTCGTGATGATAGAATGCAG 771526

RESULT 9
US-10-928-446A-193
; Sequence 193, Application US/10928446A
; Publication No. US2005027123A1
; GENERAL INFORMATION:
; APPLICANT: UNIVERSITY OF UTAH RESEARCH FOUNDATION
; TITLE OF INVENTION: VARIANTS OF NEDD4L ASSOCIATED WITH HYPERTENSION AND
; FILE REFERENCE: 0274-5785.1US
; CURRENT APPLICATION NUMBER: US/10/928,446A
; PRIOR FILING DATE: 2004-08-26
; PRIOR APPLICATION NUMBER: 60/359,741
; PRIOR FILING DATE: 2002-02-26
; NUMBER OF SEQ ID NOS: 202
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 193
; LENGTH: 1080000
; TYPE: DNA
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (999562)..(999608)
; FEATURE:
; OTHER INFORMATION: full length exon 7 range is 999547-999608
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1000354)..(1000456)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1002118)..(1002284)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1006117)..(1006249)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1007860)..(1008036)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1010940)..(1011014)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1018160)..(1018291)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1018800)..(1018919)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1020028)..(1020225)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1026659)..(1026736)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1028113)..(1028167)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1034316)..(1034374)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1041390)..(1041455)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1043121)..(1043350)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1044868)..(1044989)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1047519)..(1047589)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1050296)..(1050391)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1060368)..(1060441)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1062648)..(1062708)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1064561)..(1064620)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1066207)..(1066314)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1067768)..(1067864)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1068609)..(1068681)
; OTHER INFORMATION: exon
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1073289)..(1073388)
; OTHER INFORMATION: full length exon 30 range is 1073289-1075279
```

## US-10-928-446A-193

Query Match 22.5%; Score 29; DB 8; Length 1080000;  
Best Local Similarity 55.4%; Pred. No. 12;  
Matches 56; Conservative 0; Mismatches 45; Indels 0; Gaps 0;

Qy 16 AGCGATTACTTCGAGCATTTACTGACGACAAAGACCCGACGAGATGGTCGGGCTCTTTT 75  
Db 771426 AGCAGTTCACCTGAGACTTTCTCAGGAAAGAGTGCGATACAGTTGGTGTAGTCTTAT 771485

Qy 76 TGTGTGGTCTGTGAGCTGTGTGTCACCAACCGTATTATTCG 116  
Db 771486 ATTTGCTACGTCTCATGTGTGTCTGATGATAGATGCAG 771526

## RESULT 10

US-10-928-446A-195  
; Sequence 195, Application US/10928446A  
; Publication No. US2005027123A1

## ; GENERAL INFORMATION:

; APPLICANT: UNIVERSITY OF UTAH RESEARCH FOUNDATION  
; TITLE OF INVENTION: VARIANTS OF NEDD4L ASSOCIATED WITH HYPERTENSION AND  
; FILE REFERENCE: 0274-5785.1US  
; CURRENT APPLICATION NUMBER: US/10/928,446A  
; CURRENT FILING DATE: 2004-08-26  
; PRIOR APPLICATION NUMBER: 60/359,741  
; PRIOR FILING DATE: 2002-02-26  
; NUMBER OF SEQ ID NOS: 202  
; SOFTWARE: PatentIn Ver. 2.1  
; SEQ ID NO 195

; TYPE: DNA

; ORGANISM: Homo sapiens

; FEATURE:

; NAME/KEY: CDS

; LOCATION: (999562)..(999608)

; FEATURE:

; OTHER INFORMATION: full length exon 7 range is 999547-999608

; FEATURE:

; NAME/KEY: CDS

; LOCATION: (1000354)..(1000456)

; FEATURE:

; NAME/KEY: CDS

; LOCATION: (1002118)..(1002284)

; FEATURE:

; NAME/KEY: CDS

; LOCATION: (1006117)..(1006249)

; FEATURE:

; NAME/KEY: CDS

; LOCATION: (1007860)..(1008036)

; FEATURE:

; NAME/KEY: CDS

; LOCATION: (1010940)..(1011014)

; FEATURE:

; NAME/KEY: CDS

; LOCATION: (1018160)..(1018291)

; FEATURE:

; NAME/KEY: CDS

; LOCATION: (1018800)..(1018919)

; FEATURE:

; NAME/KEY: CDS

; LOCATION: (1020028)..(1020225)

; FEATURE:

; NAME/KEY: CDS

; LOCATION: (1026659)..(1026736)

; FEATURE:

; NAME/KEY: CDS

; LOCATION: (1028113)..(1028167)

; FEATURE:

; NAME/KEY: CDS

; LOCATION: (1034316)..(1034374)

; FEATURE:

; NAME/KEY: CDS  
; LOCATION: (1041390)..(1041455)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1043121)..(1043350)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1044868)..(1044989)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1047519)..(1047589)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1050296)..(1050391)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1060368)..(1060441)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1062648)..(1062708)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1064561)..(1064620)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1066207)..(1066314)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1067768)..(1067864)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1068609)..(1068681)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1073289)..(1073388)  
; FEATURE:

; OTHER INFORMATION: full length exon 30 range is 1073289-1075279

US-10-928-446A-195

Query Match 22.5%; Score 29; DB 8; Length 1080000;

Best Local Similarity 55.4%; Pred. No. 12;  
Matches 56; Conservative 0; Mismatches 45; Indels 0; Gaps 0;

Qy 16 AGCGATTACTTCGAGCATTTACTGACGACAAAGACCCGACGAGATGGTCGGGCTCTTTT 75  
Db 771426 AGCAGTTCACCTGAGACTTTCTCAGGAAAGAGTGCGATACAGTTGGTGTAGTCTTAT 771485

Qy 76 TGTGTGGTCTGTGAGCTGTGTGTCACCAACCGTATTATTCG 116  
Db 771486 ATTTGCTACGTCTCATGTGTGTCTGATGATAGATGCAG 771526

## RESULT 11

US-10-928-446A-197  
; Sequence 197, Application US/10928446A  
; Publication No. US2005027123A1

## ; GENERAL INFORMATION:

; APPLICANT: UNIVERSITY OF UTAH RESEARCH FOUNDATION  
; TITLE OF INVENTION: VARIANTS OF NEDD4L ASSOCIATED WITH HYPERTENSION AND  
; FILE REFERENCE: 0274-5785.1US  
; CURRENT APPLICATION NUMBER: US/10/928,446A  
; CURRENT FILING DATE: 2004-08-26  
; PRIOR APPLICATION NUMBER: 60/359,741  
; PRIOR FILING DATE: 2002-02-26  
; NUMBER OF SEQ ID NOS: 202  
; SOFTWARE: PatentIn Ver. 2.1  
; SEQ ID NO 197

; LENGTH: 1080000

; TYPE: DNA

; ORGANISM: Homo sapiens

; FEATURE:

; NAME/KEY: CDS

LOCATION: (999562)..(999608)  
OTHER INFORMATION: full length exon 7 range is 999547-999608  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1000354)..(1000456)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1002118)..(1002284)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1006117)..(1006249)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1007860)..(1008036)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1010940)..(1011014)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1018160)..(1018291)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1018800)..(1018919)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1020028)..(1020225)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1026659)..(1026736)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1028113)..(1028167)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1034316)..(1034374)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1041390)..(1041455)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1043121)..(1043350)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1044868)..(1044989)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1047519)..(1047589)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1050296)..(1050391)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1060368)..(1060441)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1062648)..(1062708)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1064561)..(1064620)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1066207)..(1066314)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1067768)..(1067864)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1068609)..(1068681)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1073289)..(1073388)  
FEATURE:

OTHER INFORMATION: full length exon 30 range is 1073289-1075279  
US-10-928-446A-197  
Query Match 22.5%; Score 29; DB 8; Length 1080000;  
Best Local Similarity 55.4%; Pred.No. 12;  
Matches 56; Conservative 0; Mismatches 45; Indels 0; Gaps 0;  
QY 16 AGCGATTACTTCGAGCATCTACTGACGACAAAGACCCGACGAGATGGTCGGGGTCTTTT 75  
DB 771426 AGCAGTTCACTTGAGACTTTCTCACGMAAAGATGCGATACAGTTGGTGTAGTCTTAT 771485  
QY 76 TGTGTGTGCTGCTGACGCTGTGTCACCGTATTATTCG 116  
DB 771486 ATTTGCTACGTGCTCATGTGTGTCGATGATAGATGCAG 771526  
RESULT 12  
US-10-928-446A-199  
Sequence 199, Application US/10928446A  
Publication No. US2005027123A1  
GENERAL INFORMATION:  
APPLICANT: UNIVERSITY OF UTAH RESEARCH FOUNDATION  
TITLE OF INVENTION: VARIANTS OF NEDD4L ASSOCIATED WITH HYPERTENSION AND  
TITLE OF INVENTION: VIRAL BUDDING  
FILE REFERENCE: 0274-5785.1US  
CURRENT APPLICATION NUMBER: US/10/928,446A  
CURRENT FILING DATE: 2004-08-26  
PRIOR APPLICATION NUMBER: 60/359,741  
PRIOR FILING DATE: 2002-02-26  
NUMBER OF SEQ ID NOS: 202  
SOFTWARE: PatentIn Ver. 2.1  
SEQ ID NO 199  
LENGTH: 1080000  
TYPE: DNA  
ORGANISM: Homo sapiens  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (999562)..(999608)  
FEATURE:  
OTHER INFORMATION: full length exon 7 range is 999547-999608  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1000354)..(1000456)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1002118)..(1002284)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1006117)..(1006249)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1007860)..(1008036)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1010940)..(1011014)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1018160)..(1018291)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1018800)..(1018919)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1020028)..(1020225)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1026659)..(1026736)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1028113)..(1028167)  
FEATURE:  
NAME/KEY: CDS  
LOCATION: (1034316)..(1034374)  
FEATURE:

```
; NAME/KEY: allele  
; LOCATION: (827008)..(827008)  
; OTHER INFORMATION: the 'n' at position 827008 may be 'a' or 'g'  
; FEATURE:  
; NAME/KEY: 5'UTR  
; LOCATION: (872617)..(872860)  
; OTHER INFORMATION: Exon 2a  
; FEATURE:  
; NAME/KEY: 5'UTR  
; LOCATION: (904952)..(905030)  
; OTHER INFORMATION: Exon 2d  
; FEATURE:  
; NAME/KEY: 5'UTR  
; LOCATION: (922549)..(922630)  
; OTHER INFORMATION: Exon 3  
; FEATURE:  
; NAME/KEY: 5'UTR  
; LOCATION: (926021)..(926059)  
; FEATURE:  
; NAME/KEY: 5'UTR  
; LOCATION: (929123)..(929176)  
; OTHER INFORMATION: Exon 5  
; FEATURE:  
; NAME/KEY: 5'UTR  
; LOCATION: (993104)..(993154)  
; OTHER INFORMATION: Exon 6  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (999562)..(999608)  
; OTHER INFORMATION: Full exon range is 999547-999608  
; FEATURE:  
; NAME/KEY: exon  
; LOCATION: (1000354)..(1000456)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1000354)..(1000456)  
; FEATURE:  
; NAME/KEY: exon  
; LOCATION: (1002118)..(1002284)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1002118)..(1002284)  
; FEATURE:  
; NAME/KEY: exon  
; LOCATION: (1006117)..(1006249)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1006117)..(1006249)  
; FEATURE:  
; NAME/KEY: exon  
; LOCATION: (1007860)..(1008036)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1007860)..(1008036)  
; FEATURE:  
; NAME/KEY: exon  
; LOCATION: (1010940)..(1011014)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1010940)..(1011014)  
; FEATURE:  
; NAME/KEY: exon  
; LOCATION: (1018160)..(1018291)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1018160)..(1018291)  
; FEATURE:  
; NAME/KEY: exon  
; LOCATION: (1018900)..(1018919)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1018900)..(1018919)  
; FEATURE:
```

---

```
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1041390)..(1041455)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1043121)..(1043350)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1044868)..(1044989)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1047519)..(1047589)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1050296)..(1050391)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1060368)..(1060441)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1062648)..(1062708)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1064561)..(1064620)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1066207)..(1066314)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1067768)..(1067864)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1068609)..(1068681)  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (1073289)..(1073388)  
; FEATURE:  
; OTHER INFORMATION: full length exon 30 range is 1073289-1075279
```

US-10-928-446A-199

```
Query Match          22.5%; Score 29; DB 8; Length 1080000;  
Best Local Similarity 55.4%; Pred. No. 12;  
Matches 56; Conservative 0; Mismatches 45; Indels 0; Gaps 0;
```

		Qy	16	AGCGATTACTTCGACGATTACTCAGCAACAAGACC CGCACGGAGTGTCCGGGCTTTT	75
Db	771426	AGCAGITCATTTGAGACTTCTCACGGAAGAAGTGCGTAGAGTTGGTGCAGTACTTAT	771485	           	
Qy	76	TGTTGGTGCTGTGACGTTGTGCCAACCGTATTATTCGG	116	           	
Db	771486	ATTTCGTACGTGCTCATGTCTGCTGATGATAAATGCAG	771526	           	

RESULT 13

US-10-928-446A-201

```
; Sequence 201, Application US/10928446A  
; Publication No. US2005027123A1  
GENERAL INFORMATION:  
APPLICANT: UNIVERSITY OF UTAH RESEARCH FOUNDATION  
TITLE OF INVENTION: VARIANTS OF NEDD4L ASSOCIATED WITH HYPERTENSION AND TITLE OF INVENTION: VIRAL BUDDING  
FILE REFERENCE: 0274-5785.IUS  
CURRENT APPLICATION NUMBER: US/10/928,446A  
PRIOR FILING DATE: 2004-08-26  
PRIORITY CLAIMS: PENDING  
NUMBER OF SEQ ID NOS: 202  
SOFTWARE: Patentin Ver. 2.1  
SEQ ID NO 201  
LENGTH: 1080000  
TYPE: DNA  
ORGANISM: Homo sapiens  
FEATURE:
```

```
; NAME/KEY: exon
; LOCATION: (1020028) .. (1020225)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1020028) .. (1020225)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1026659) .. (1026736)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1026659) .. (1026736)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1028113) .. (1028167)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1028113) .. (1028167)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1034316) .. (1034374)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1034316) .. (1034374)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1041390) .. (1041455)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1041390) .. (1041455)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1043121) .. (1043350)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1043121) .. (1043350)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1044868) .. (1044989)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1044868) .. (1044989)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1047519) .. (1047589)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1047519) .. (1047589)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1050296) .. (1050391)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1050296) .. (1050391)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1060368) .. (1060441)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1060368) .. (1060441)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1062648) .. (1062708)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1062648) .. (1062708)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1064561) .. (1064620)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1064561) .. (1064620)
; FEATURE:
; NAME/KEY: exon
```

```
; LOCATION: (1066207) .. (1066314)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1066207) .. (1066314)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1067768) .. (1067864)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1067768) .. (1067864)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1068609) .. (1068681)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1068609) .. (1068681)
; FEATURE:
; NAME/KEY: exon
; LOCATION: (1073289) .. (1075279)
; FEATURE:
; NAME/KEY: CDS
; LOCATION: (1073289) .. (1073388)
; US-10-928-446A-201
```

Query Match 22.5%; Score 29; DB 8; Length 1080000;  
Best Local Similarity 55.4%; Pred. No. 12;  
Matches 56; Conservative 0; Mismatches 45; Indels 0; Gaps 0;

```
Qy 16 AGCGATTACTTCGAGCATTACTGACGACAAAGACCCGACCGAGATGGTCGGGCTCTTTT 75
Db 771426 AGCAGTTCAGTCTGAGACTTCTCACGGAAGAAGTCGATACAGTTGGTGTAGTCTTAT 771485

Qy 76 TGTGTGGTGTGCTGACGTGTTGTCCAAACCGTATTATTCCG 116
Db 771486 ATTTGCTACGTGCTCATGTGTTGCTGATGATAGATGCAG 771526
```

## RESULT 14

```
US-11-124-368A-2924/c
; Sequence 2924, Application US/11124368A
; Publication No. US20050287559A1
; GENERAL INFORMATION:
; APPLICANT: Michele Cargill
; APPLICANT: James J. Devlin
; TITLE OF INVENTION: Genetic Polymorphisms Associated with
; TITLE OF INVENTION: Vascular Diseases, Methods of Detection and Uses Thereof
; FILE REFERENCE: CL001524
; CURRENT APPLICATION NUMBER: US/11/124,368A
; CURRENT FILING DATE: 2005-05-09
; PRIOR APPLICATION NUMBER: US 60/568,845
; PRIOR FILING DATE: 2004-05-07
; PRIOR APPLICATION NUMBER: US 60/625,936
; PRIOR FILING DATE: 2004-11-09
; NUMBER OF SEQ ID NOS: 21112
; SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO 2924
; LENGTH: 79134
; TYPE: DNA
; ORGANISM: Homo sapiens
; US-11-124-368A-2924
```

Query Match 22.3%; Score 28.8; DB 12; Length 79134;  
Best Local Similarity 58.0%; Pred. No. 7;  
Matches 51; Conservative 0; Mismatches 37; Indels 0; Gaps 0;

```
Qy 23 ACTTCGAGCATTACTGACGACAAAGACCCGACCGAGATGGTCGGGCTCTTTTGTG 82
Db 76157 AGTTCTAGGATTACAGACATGAGACACACGCGCGCTGATATATAACTTTTTTTT 76098

Qy 83 GTGCTGTGACGTGTGTCCTCCAAACCGTATT 110
Db 76097 TTTTGTGATGAAGTCTCACTCTGTT 76070
```



GenCore version 5.1.7  
Copyright (c) 1993 - 2006 Bioceleration Ltd.  
OM nucleic - nucleic search, using sw model  
Run on: March 7, 2006, 12:14:00 ; Search time 354.047 Seconds  
(without alignments)  
3013.020 Million cell updates/sec

Title: US-10-782-899-1\_COPY\_472\_600  
Perfect score: 129  
Sequence: 1 tgaccttcgtcgtagcga.....tattccggactgttcagcg 129

Scoring table: IDENTITY NUC  
Gapop 10.0 , Gapext 1.0

Searched: 9793542 seqs, 4134689005 residues  
Total number of hits satisfying chosen parameters: 19587084

Minimum DB seq length: 0  
Maximum DB seq length: 2000000000  
Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database : Published Applications\_NA\_Main:\*

- 1: /cgn2\_6/ptodata/1/pubpna/US07\_PUBCOMB.seq:\*
- 2: /cgn2\_6/ptodata/1/pubpna/US08\_PUBCOMB.seq:\*
- 3: /cgn2\_6/ptodata/1/pubpna/US09A\_PUBCOMB.seq:\*
- 4: /cgn2\_6/ptodata/1/pubpna/US09B\_PUBCOMB.seq:\*
- 5: /cgn2\_6/ptodata/1/pubpna/US10A\_PUBCOMB.seq:\*
- 6: /cgn2\_6/ptodata/1/pubpna/US10B\_PUBCOMB.seq:\*
- 7: /cgn2\_6/ptodata/1/pubpna/US10C\_PUBCOMB.seq:\*
- 8: /cgn2\_6/ptodata/1/pubpna/US10D\_PUBCOMB.seq:\*
- 9: /cgn2\_6/ptodata/1/pubpna/US10E\_PUBCOMB.seq:\*
- 10: /cgn2\_6/ptodata/1/pubpna/US11\_PUBCOMB.seq:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	129	100.0	600	3	US-09-816-391A-1
2	129	100.0	600	8	US-10-782-899-1
3	129	100.0	2256646	7	US-10-470-565-1
4	29.4	22.8	885	9	US-10-617-320-306
5	29.4	22.8	1377	7	US-10-687-677-1
6	29.4	22.8	3520	3	US-09-814-353-19115
7	29.2	22.6	634	7	US-10-332-859-15
8	29.2	22.6	1013	7	US-10-332-859-232
9	29.2	22.6	1836	7	US-10-332-859-17
10	29	22.5	344805	8	US-10-779-271-1
11	29	22.5	354592	9	US-10-737-082-70
12	29	22.5	354592	9	US-10-765-790-70
13	28.8	22.3	766	7	US-10-332-859-317
14	28.6	22.2	1043	8	US-10-425-115-152385
15	28.4	22.0	936	3	US-09-503-410-31
16	28.4	22.0	936	5	US-10-027-805-31
17	28.4	22.0	936	5	US-10-027-804-31
18	28.4	22.0	2451	3	US-09-938-842A-64
19	28.4	22.0	2451	3	US-09-938-842A-64
20	28.2	21.9	599	6	US-10-029-386-13097
21	28.2	21.9	1152	9	US-10-450-763-9386
22	28.2	21.9	3519	7	US-10-437-963-80607
23	28	21.7	165221	5	US-10-087-192-1015

C 24	27.8	21.6	483	7	US-10-424-599-128897	Sequence 128897,
C 25	27.8	21.6	2037	5	US-10-052-586-591	Sequence 591, App
C 26	27.8	21.6	2037	5	US-10-174-590-591	Sequence 591, App
C 27	27.8	21.6	2037	5	US-10-176-758-591	Sequence 591, App
C 28	27.8	21.6	2037	5	US-10-175-737-591	Sequence 591, App
C 29	27.8	21.6	2037	5	US-10-174-581-591	Sequence 591, App
C 30	27.8	21.6	2037	5	US-10-176-483-591	Sequence 591, App
C 31	27.8	21.6	2037	5	US-10-176-749-591	Sequence 591, App
C 32	27.8	21.6	2037	5	US-10-176-914-591	Sequence 591, App
C 33	27.8	21.6	2037	5	US-10-176-915-591	Sequence 591, App
C 34	27.8	21.6	2037	5	US-10-173-706-591	Sequence 591, App
C 35	27.8	21.6	2037	5	US-10-175-738-591	Sequence 591, App
C 36	27.8	21.6	2037	5	US-10-175-738-591	Sequence 591, App
C 37	27.8	21.6	2037	5	US-10-176-482-591	Sequence 591, App
C 38	27.8	21.6	2037	5	US-10-176-757-591	Sequence 591, App
C 39	27.8	21.6	2037	5	US-10-176-913-591	Sequence 591, App
C 40	27.8	21.6	2037	5	US-10-180-552-591	Sequence 591, App
C 41	27.8	21.6	2037	5	US-10-180-557-591	Sequence 591, App
C 42	27.8	21.6	2037	5	US-10-173-700-591	Sequence 591, App
C 43	27.8	21.6	2037	5	US-10-174-572-591	Sequence 591, App
C 44	27.8	21.6	2037	5	US-10-174-579-591	Sequence 591, App
C 45	27.8	21.6	2037	5	US-10-174-582-591	Sequence 591, App

ALIGNMENTS

RESULT 1  
US-09-816-391A-1  
; Sequence 1, Application US/09816391A  
; Patent No. US2002005486SA1  
; GENERAL INFORMATION:  
; APPLICANT: FUJIMORI, Minoru  
; APPLICANT: TANIGUCHI, Shunichiro  
; APPLICANT: AMANO, Jun  
; APPLICANT: YAZAWA, Kazuyuki  
; APPLICANT: KANO, Yasunobu  
; APPLICANT: NAKAMURA, Toshiyuki  
; APPLICANT: SASAKI, Takayuki  
; TITLE OF INVENTION: Anaerobic bacterium as a drug for cancer gene therapy  
; FILE REFERENCE: 2001-WMC/01736  
; CURRENT APPLICATION NUMBER: US/09/816,391A  
; CURRENT FILING DATE: 2001-03-26  
; PRIOR APPLICATION NUMBER: JP 00/287688  
; PRIOR FILING DATE: 2000-09-21  
; NUMBER OF SEQ ID NOS: 3  
; SEQ ID NO 1  
; LENGTH: 600  
; TYPE: DNA  
; ORGANISM: Bifidobacterium longum  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (193)...(471)  
US-09-816-391A-1

Query Match 100.0%; Score 129; DB 3; Length 600;  
Best Local Similarity 100.0%; Pred. No. 5e-37;  
Matches 129; Conservative 0; Mismatches 0; Gaps 0;  
QY 1 TGACCTTCGCTGCTAGCGATTACTTCGAGCATTTACTGACGACAAAGACCCCGACCGAGA 60  
Db 472 TGACCTTCGCTGCTAGCGATTACTTCGAGCATTTACTGACGACAAAGACCCCGACCGAGA 531  
QY 61 TGGTCGGGGCTCTTTTGTGTGCTGTGAGCTGTGCTCCACCGCTATTATTCGCGACT 120  
Db 532 TGGTCGGGGCTCTTTTGTGTGCTGTGAGCTGTGCTCCACCGCTATTATTCGCGACT 591  
QY 121 AGTTTCAGCG 129  
Db 592 AGTTTCAGCG 600

RESULT 2

US-10-782-899-1  
; Sequence 1, Application US/10782899  
; Publication No. US20050025745A1  
; GENERAL INFORMATION:  
; APPLICANT: FUJIMORI, MINORU  
; APPLICANT: TANIGUCHI, SHUNICHIRO  
; APPLICANT: ANANO, JUN  
; APPLICANT: YAZAWA, KAZUYUKI  
; APPLICANT: KANO, YASUNOBU  
; APPLICANT: NAKAMURA, TOSHIYUKI  
; APPLICANT: SASAKI, TAKAYUKI  
; TITLE OF INVENTION: ANAEROBIC BACTERIUM AS A DRUG FOR CANCER GENE THERAPY  
; FILE REFERENCE: 671308-2001.1  
; CURRENT APPLICATION NUMBER: US/10/782,899  
; CURRENT FILING DATE: 2004-02-23  
; PRIOR APPLICATION NUMBER: 09/816,391  
; PRIOR FILING DATE: 2001-03-26  
; PRIOR APPLICATION NUMBER: JP 2000-287688  
; PRIOR FILING DATE: 2000-09-21  
; NUMBER OF SEQ ID NOS: 4  
; SOFTWARE: PatentIn ver. 3.2  
; SEQ ID NO 1  
; LENGTH: 600  
; TYPE: DNA  
; ORGANISM: Bifidobacterium longum  
; FEATURE:  
; NAME/KEY: CDS  
; LOCATION: (193)..(471)  
US-10-782-899-1  
Query Match 100.0%; Score 129; DB 8; Length 600;  
Best Local Similarity 100.0%; Pred. No. 5e-37;  
Matches 129; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 1 TGACCTTCTGCTAGCGATTACTTCGAGCATTACTGACGACAAAGACCCGACCGAGA 60  
DB 472 TGACCTTCTGCTAGCGATTACTTCGAGCATTACTGACGACAAAGACCCGACCGAGA 531  
QY 61 TGGTCGGGGTCTTTTGTGCTGCTGACGCTGTTGTCACCGATTATTATTCGGGACT 120  
DB 532 TGGTCGGGGTCTTTTGTGCTGCTGACGCTGTTGTCACCGATTATTATTCGGGACT 591  
QY 121 AGTTCAGCG 129  
DB 592 AGTTCAGCG 600  
RESULT 3  
; Sequence 1, Application US/10470565  
; Publication No. US20040126870A1  
; GENERAL INFORMATION:  
; APPLICANT: Societe des Produits Nestle S.A.  
; TITLE OF INVENTION: NCC2705 - the genome of a Bifidobacterium  
; FILE REFERENCE: 80290/WO  
; CURRENT APPLICATION NUMBER: US/10/470,565  
; CURRENT FILING DATE: 2003-07-29  
; PRIOR APPLICATION NUMBER: EP 01102050.0  
; PRIOR FILING DATE: 2001-01-30  
; NUMBER OF SEQ ID NOS: 2  
; SOFTWARE: PatentIn version 3.1  
; SEQ ID NO 1  
; LENGTH: 2256646  
; TYPE: DNA  
; ORGANISM: Bifidobacterium longum  
US-10-470-565-1  
Query Match 100.0%; Score 129; DB 7; Length 2256646;  
Best Local Similarity 100.0%; Pred. No. 6.6e-36;  
Matches 129; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 1 TGACCTTCTGCTAGCGATTACTTCGAGCATTACTGACGACAAAGACCCGACCGAGA 60

DB 2241699 TGACCTTCTGCTAGCGATTACTTCGAGCATTACTGACGACAAAGACCCGACCGAGA 2241640  
QY 61 TGGTCGGGGTCTTTTGTGCTGCTGACGCTGTTGTCACCGATTATTATTCGGGACT 120  
DB 2241639 TGGTCGGGGTCTTTTGTGCTGCTGACGCTGTTGTCACCGATTATTATTCGGGACT 2241580  
QY 121 AGTTCAGCG 129  
DB 2241579 AGTTCAGCG 2241571  
RESULT 4  
US-10-617-320-306  
; Sequence 306, Application US/10617320  
; Publication No. US20050136404A1  
; GENERAL INFORMATION:  
; APPLICANT: Lythn A Doucette-Stamm and David Bush  
; TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID  
; SEQUENCES RELATING TO STREPTOCOCCUS PNEUMONIAE  
; THERAPEUTICS  
; NUMBER OF SEQUENCES: 5206  
; CORRESPONDENCE ADDRESS:  
; ADDRESSEE: GENOME THERAPEUTICS CORPORATION  
; STREET: 100 Beaver Street  
; CITY: Waltham  
; STATE: Massachusetts  
; COUNTRY: USA  
; ZIP: 02354  
; COMPUTER READABLE FORM:  
; MEDIUM TYPE: CD-ROM ISO9660  
; COMPUTER: <Unknown>  
; OPERATING SYSTEM: <Unknown>  
; SOFTWARE: <Unknown>  
; CURRENT APPLICATION DATA:  
; APPLICATION NUMBER: US/10/617,320  
; FILING DATE: 10-Jul-2003  
; PRIOR APPLICATION DATA:  
; APPLICATION NUMBER: US/09/107,433  
; FILING DATE: 30-Jun-1998  
; APPLICATION NUMBER: 60/ 085131  
; FILING DATE: May 12, 1998  
; APPLICATION NUMBER: 60/051553  
; FILING DATE: July 2, 1997  
; ATTORNEY/AGENT INFORMATION:  
; NAME: Ariniello, Pamela Deneke  
; REGISTRATION NUMBER: 40,489  
; REFERENCE/DOCKET NUMBER: GTC-011  
; TELECOMMUNICATION INFORMATION:  
; TELEPHONE: (781)893-5007  
; TELEFAX: (781)893-8277  
; INFORMATION FOR SEQ ID NO: 306:  
; SEQUENCE CHARACTERISTICS:  
; LENGTH: 885 base pairs  
; TYPE: nucleic acid  
; STRANDEDNESS: double  
; TOPOLOGY: circular  
; MOLECULE TYPE: DNA (genomic)  
; HYPOTHETICAL: NO  
; ANTI-SENSE: NO  
; ORIGINAL SOURCE:  
; ORGANISM: Streptococcus pneumoniae  
; FEATURE:  
; NAME/KEY: misc\_feature  
; LOCATION: (B) LOCATION 1...885  
; SEQUENCE DESCRIPTION: SEQ ID NO: 306:  
US-10-617-320-306  
Query Match 22.8%; Score 29.4; DB 9; Length 885;  
Best Local Similarity 58.8%; Pred. No. 3.5;  
Matches 51; Conservative 0; Mismatches 36; Indels 0; Gaps 0;  
QY 32 ATTACTGACGACAAAGACCCGACCGAGATGGTCGGGGTCTTTTGTGTTGGTGTGTA 91



Db 604 ATTACAGATATCAACAGGCTTACCTAAATCGTGTGACCTTTCTGTTGAGCTGATGGGG 663

QY 92 CGTGTGTTGCCAACCGGTATTATCCGGA 118

Db 664 CGTGGTTTGGCTGGTTGGATACGGA 690

RESULT 5

US-10-687-677-1/c

Sequence 1, Application US/10687677

Publication No. US20040142419A1

GENERAL INFORMATION:

APPLICANT: Guy, John

TITLE OF INVENTION: REDUCING CELLULAR DYSFUNCTION CAUSED BY MITOCHONDRIAL GENE

FILE OF INVENTION: MUTATIONS

FILE REFERENCE: 5853-324

CURRENT APPLICATION NUMBER: US/10/687,677

CURRENT FILING DATE: 2003-10-17

NUMBER OF SEQ ID NOS: 2

SOFTWARE: Patent in version 3.2

SEQ ID NO 1

LENGTH: 1377

TYPE: DNA

ORGANISM: Homo sapiens

US-10-687-677-1

Query Match 22.8%; Score 29.4; DB 7; Length 1377;

Best Local Similarity 63.4%; Pred. No. 4;

Matches 45; Conservative 0; Mismatches 26; Indels 0; Gaps 0;

QY 57 GAGATGGTCGGGCTCTTTTGTGTCGTGAGCGTGTGTCACCGGTATATATCCG 116

Db 1292 GTCAAGCTGGCTTCATGTTGTGATGTGTGTCAGGCTGCCCACTGTGTGTG 1233

QY 117 GACTAGTTTCAG 127

Db 1232 AACATGTACAG 1222

RESULT 6

US-09-814-353-19115

Sequence 19115, Application US/09814353

Publication No. US20030165831A1

GENERAL INFORMATION:

APPLICANT: Lee, John

APPLICANT: Thompson, Pamela

APPLICANT: Lillie, James

TITLE OF INVENTION: NOVEL GENES, COMPOSITIONS, KITS, AND METHODS FOR

TITLE OF INVENTION: IDENTIFICATION, ASSESSMENT, PREVENTION, AND

TITLE OF INVENTION: THERAPY OF OVARIAN CANCER

FILE REFERENCE: MRI-006B

CURRENT APPLICATION NUMBER: US/09/814,353

CURRENT FILING DATE: 2001-03-21

PRIOR APPLICATION NUMBER: US 60/191,031

PRIOR FILING DATE: 2000-03-21

PRIOR APPLICATION NUMBER: US 60/207,124

PRIOR FILING DATE: 2000-05-25

PRIOR APPLICATION NUMBER: US 60/211,940

PRIOR FILING DATE: 2000-06-15

PRIOR APPLICATION NUMBER: US 60/216,820

PRIOR FILING DATE: 2000-07-07

PRIOR APPLICATION NUMBER: US 60/220,661

PRIOR FILING DATE: 2000-07-25

PRIOR APPLICATION NUMBER: US 60/257,672

PRIOR FILING DATE: 2000-12-21

NUMBER OF SEQ ID NOS: 22037

SOFTWARE: FastSeq for Windows Version 4.0

SEQ ID NO 19115

LENGTH: 3520

TYPE: DNA

ORGANISM: Homo sapiens

US-09-814-353-19115

Query Match 22.8%; Score 29.4; DB 3; Length 3520;

Best Local Similarity 54.1%; Pred. No. 5.4;

Matches 60; Conservative 0; Mismatches 51; Indels 0; Gaps 0;

QY 4 CCTTCGCTCGTAGCGATTACTTCGAGCATTACTGACGACAGACACCCGACGAGATGG 63

Db 27 CCTCTACTGCTCATCTCTGCTTACCTGTGTTTTTTAGTGATTAATCCCTCACCTGTGG 86

QY 64 TCGGGGTCTTTTTTGTGTCGTGTCGTGAGCGTGTGTCACACCGTATTATTTC 114

Db 87 GCGATGTGTTTTTTTGTGGGGCTGTAACTGCTGGCCCACTTCATAATCC 137

RESULT 7

US-10-332-859-15/c

Sequence 15, Application US/10332859

Publication No. US20040088746A1

GENERAL INFORMATION:

APPLICANT: Grimm, Stefan

APPLICANT: Schoenfeld, Nicole

APPLICANT: Braziulis, Erik

APPLICANT: Cramer, Ursula

APPLICANT: Gewies, Andreas

APPLICANT: Voss, Frank

APPLICANT: Mund, Thomas

APPLICANT: Albayrak, Timur

APPLICANT: Gille, Hendrik

APPLICANT: Klein, Matthias

APPLICANT: Bauer, Manuel

TITLE OF INVENTION: Apoptosis-Inducing DNA Sequences

FILE REFERENCE: 2923-0133

CURRENT APPLICATION NUMBER: US/10/332,859

CURRENT FILING DATE: 2003-01-14

PRIOR APPLICATION NUMBER: PCT/EP01/08170

PRIOR FILING DATE: 2001-07-13

NUMBER OF SEQ ID NOS: 355

SOFTWARE: Patent in version 3.2

SEQ ID NO 15

LENGTH: 634

TYPE: DNA

ORGANISM: Mus musculus

US-10-332-859-15

Query Match 22.6%; Score 29.2; DB 7; Length 634;

Best Local Similarity 59.8%; Pred. No. 3.8;

Matches 49; Conservative 0; Mismatches 33; Indels 0; Gaps 0;

QY 48 ACCCGACCGAGATGTCGGGGCTCTTTTGTGTCGTGACCGTGTGTCACACCGT 107

Db 245 ATCCCTCACCTCTCGGGCGATGTGTTTCTTTGTGGGCTGTAACTGCTGGCCACTTC 186

QY 108 ATTATTCCGGACTAGTTTCAGCG 129

Db 185 ATCAATGCCCTACTTGTGGACG 164

RESULT 8

US-10-332-859-232

Sequence 232, Application US/10332859

Publication No. US20040088746A1

GENERAL INFORMATION:

APPLICANT: Grimm, Stefan

APPLICANT: Schoenfeld, Nicole

APPLICANT: Braziulis, Erik

APPLICANT: Cramer, Ursula

APPLICANT: Gewies, Andreas

APPLICANT: Voss, Frank

APPLICANT: Mund, Thomas

APPLICANT: Albayrak, Timur

APPLICANT: Gille, Hendrik

APPLICANT: Klein, Matthias

APPLICANT: Bauer, Manuel

TITLE OF INVENTION: Apoptosis-Inducing DNA Sequences

```
; FILE REFERENCE: 2923-0133
; CURRENT APPLICATION NUMBER: US/10/332,859
; CURRENT FILING DATE: 2003-01-14
; PRIOR APPLICATION NUMBER: PCT/EP01/08170
; PRIOR FILING DATE: 2001-07-13
; NUMBER OF SEQ ID NOS: 355
; SOFTWARE: PatentIn version 3.2
; SEQ ID NO 232
; LENGTH: 1013
; TYPE: DNA
; ORGANISM: Mus musculus
US-10-332-859-232

Query Match      22.6%; Score 29.2; DB 7; Length 1013;
Best Local Similarity 59.8%; Pred. No. 4.3;
Matches 49; Conservative 0; Mismatches 33; Indels 0; Gaps 0;

Qy 48 ACCCGACCGAGATGTCGGGGCTTTTGTGTGGTGTGTGACGTGTGTGTCACACCGT 107
    |||||
Db 769 ATCCCTCACCTCTCGGGCGATGTGGTTTCTTGTGGGGCTGTAACTGCTGGCCCACTTC 828

Qy 108 ATTATTCGGAGTAGTTTCAGCG 129
    |||||
Db 829 ATCAATGCCTACTTGGTGGACG 850

RESULT 9
US-10-332-859-17
; Sequence 17, Application US/10332859
; Publication No. US20040088746A1
; GENERAL INFORMATION:
; APPLICANT: Grimm, Stefan
; APPLICANT: Schoenfeld, Nicole
; APPLICANT: Brazilius, Erik
; APPLICANT: Cramer, Ursula
; APPLICANT: Gewies, Andreas
; APPLICANT: Voss, Frank
; APPLICANT: Mund, Thomas
; APPLICANT: Albayrak, Timur
; APPLICANT: Gille, Hendrik
; APPLICANT: Klein, Matthias
; APPLICANT: Bauer, Manuel
; TITLE OF INVENTION: Apoptosis-Inducing DNA Sequences
; FILE REFERENCE: 2923-0133
; CURRENT APPLICATION NUMBER: US/10/332,859
; CURRENT FILING DATE: 2003-01-14
; PRIOR APPLICATION NUMBER: PCT/EP01/08170
; PRIOR FILING DATE: 2001-07-13
; NUMBER OF SEQ ID NOS: 355
; SOFTWARE: PatentIn version 3.2
; SEQ ID NO 17
; LENGTH: 1836
; TYPE: DNA
; ORGANISM: Mus musculus
US-10-332-859-17

Query Match      22.6%; Score 29.2; DB 7; Length 1836;
Best Local Similarity 59.8%; Pred. No. 5.2;
Matches 49; Conservative 0; Mismatches 33; Indels 0; Gaps 0;

Qy 48 ACCCGACCGAGATGTCGGGGCTTTTGTGTGGTGTGTGACGTGTGTGTCACACCGT 107
    |||||
Db 813 ATCCCTCACCTCTCGGGCGATGTGGTTTCTTGTGGGGCTGTAACTGCTGGCCCACTTC 872

Qy 108 ATTATTCGGAGTAGTTTCAGCG 129
    |||||
Db 873 ATCAATGCCTACTTGGTGGACG 894

RESULT 10
US-10-779-271-1
; Sequence 1, Application US/10779271
; Publication No. US20040220387A1
```

```
; GENERAL INFORMATION:
; APPLICANT: AstraZeneca AB
; TITLE OF INVENTION: Methods
; FILE REFERENCE: ASZD-P02-251
; CURRENT APPLICATION NUMBER: US/10/779,271
; CURRENT FILING DATE: 2004-02-13
; PRIOR APPLICATION NUMBER: 09/463,844
; PRIOR FILING DATE: 2000-02-01
; PRIOR APPLICATION NUMBER: PCT/GB98/00259
; PRIOR FILING DATE: 1998-07-28
; PRIOR APPLICATION NUMBER: 9716162.4
; PRIOR FILING DATE: 1997-08-97
; PRIOR APPLICATION NUMBER: 60/535,986
; PRIOR FILING DATE: 2004-01-12
; NUMBER OF SEQ ID NOS: 29
; SOFTWARE: PatentIn version 3.2
; SEQ ID NO 1
; LENGTH: 344805
; TYPE: DNA
; ORGANISM: Homo sapiens
US-10-779-271-1

Query Match      22.5%; Score 29; DB 8; Length 344805;
Best Local Similarity 55.4%; Pred. No. 32;
Matches 56; Conservative 0; Mismatches 45; Indels 0; Gaps 0;

Qy 16 AGCGATTACTTCGAGCATTTACTGACGACAAGACCCGACCGAGATGGTCGGGTCCTTTT 75
    |||||
Db 47662 AGCAGTTCACTTGAGACTTTCTCACGGAAGAAGTGGGATACAGTTGGTGGTAGTCTTAT 47721

Qy 76 TGTGTGGTGTCTGTGACGTGTGTGTCACACCGTATTATTCCG 116
    |||||
Db 47722 ATTTGCTACGTGCTCATGTGTGTCATGATAGATGATGCAG 47762

RESULT 11
US-10-737-082-70
; Sequence 70, Application US/10737082
; Publication No. US20050130170A1
; GENERAL INFORMATION:
; APPLICANT: Bayer Healthcare LLC
; APPLICANT: Beard, Chris
; APPLICANT: Burgess, Chris
; APPLICANT: Gannon, Allison
; APPLICANT: Harvey, Jeanne
; APPLICANT: Lechner, John F.
; APPLICANT: Li, Zheng
; TITLE OF INVENTION: Identification and Verification of Methylation Marker Sequences
; FILE REFERENCE: 1657/2032
; CURRENT APPLICATION NUMBER: US/10/737,082
; CURRENT FILING DATE: 2003-12-16
; PRIOR APPLICATION NUMBER: US 10/737,082
; PRIOR FILING DATE: 2003-12-16
; NUMBER OF SEQ ID NOS: 300
; SOFTWARE: PatentIn version 3.2
; SEQ ID NO 70
; LENGTH: 354592
; TYPE: DNA
; ORGANISM: Homo sapiens
US-10-737-082-70

Query Match      22.5%; Score 29; DB 9; Length 354592;
Best Local Similarity 55.4%; Pred. No. 32;
Matches 56; Conservative 0; Mismatches 45; Indels 0; Gaps 0;

Qy 16 AGCGATTACTTCGAGCATTTACTGACGACAAGACCCGACCGAGATGGTCGGGTCCTTTT 75
    |||||
Db 50053 AGCAGTTCACTTGAGACTTTCTCACGGAAGAAGTGGGATACAGTTGGTGGTAGTCTTAT 50112

Qy 76 TGTGTGGTGTCTGTGACGTGTGTGTCACACCGTATTATTCCG 116
    |||||
Db 50113 ATTTGCTACGTGCTCATGTGTGTCATGATAGATGATGCAG 50153
```

QY 48 ACCCCGACCGAGATGGTCGGGGTCTTTTGTGTGGTGCTGTGACGTTGTGTCAACCGT 107  
| | | | : | | | | | | | | | | | | | | | |  
Db 229 ATCCCTCACCTCCTCGGGCGATGCGTTTTCTTGTGGGCTGTAACTCTGGCCACTTC 288  
| | | | | | | | | | | | | | | | | | | | |  
QY 108 ATTATCCGGACTAGTTTCAGG 129  
| | | | | | | | | | | | | | | | | | | | |  
Db 289 ATCAATGCCTACTTGGTGGACG 310  
| | | | | | | | | | | | | | | | | | | | |

RESULT 14  
US-10-425-115-152385/c  
; Sequence 152385, Application US/10425115  
; Publication No. US20040214272A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa, Thomas J.  
; APPLICANT: Kovalic, David K.  
; APPLICANT: Zhou, Yihua  
; APPLICANT: Cao, Yongwei  
; TITLE OF INVENTION: Nucleic Acid Molecules and Other Molecules Associated With  
; TITLE OF INVENTION: Plants  
; FILE REFERENCE: 38-21(53222)B  
; CURRENT APPLICATION NUMBER: US/10/425,115  
; CURRENT FILING DATE: 2003-04-28  
; NUMBER OF SEQ ID NOS: 369326  
; SEQ ID NO 152385  
; LENGTH: 1043  
; TYPE: DNA  
; ORGANISM: Zea mays  
; FEATURE:  
; OTHER INFORMATION: Clone ID: MRT4577\_70558C.1  
US-10-425-115-152385

Query Match 22.2%; Score 28.6; DB 8; Length 1043;  
Best Local Similarity 55.6%; Pred. No. 7.3;  
Matches 55; Conservative 0; Mismatches 44; Indels 0; Gaps 0;

QY 19 GATTACTCGAGCATTAAGTACGACAAGACCACCGAGATGTCGGGCTCTTTTGT 78  
| | | | | | | | | | | | | | | | | | | | |  
Db 1041 GCTGACTTCCACCTTGATGACTCAAGGAGCAGAGTAGTGGTCGGGATCTGGGTGC 982  
| | | | | | | | | | | | | | | | | | | | |

QY 79 TGTGGTGTGTGACGTGTGTGCCAACCGTATTATTCCGG 117  
| | | | | | | | | | | | | | | | | | | | |  
Db 981 TCCTGAGCATCGGGCTCTTCAGGAACCCGAACTTCCCG 943  
| | | | | | | | | | | | | | | | | | | | |

RESULT 15  
US-09-903-410-31  
; Sequence 31, Application US/09903410  
; Patent No. US20020146799A1  
; GENERAL INFORMATION:  
; APPLICANT: DIVERSA CORPORATION  
; APPLICANT: ROBERTSON, Dan  
; APPLICANT: MURPHY, Dennis  
; APPLICANT: REID, John  
; APPLICANT: MAFFIA, Anthony  
; APPLICANT: LINK, Steven  
; APPLICANT: SWANSON, Ronald  
; APPLICANT: WARREN, Patrick  
; APPLICANT: KOSMOTKA, Anna  
; TITLE OF INVENTION: ENZYMES HAVING ESTERASE ACTIVITY AND METHODS OF USE THEREOF  
; FILE REFERENCE: DIVER180-2  
; CURRENT APPLICATION NUMBER: US/09/903,410  
; CURRENT FILING DATE: 2001-07-10  
; PRIOR APPLICATION NUMBER: US 09/382,242  
; PRIOR FILING DATE: 1999-08-24  
; PRIOR APPLICATION NUMBER: US 08/602,359  
; PRIOR FILING DATE: 1996-02-16  
; NUMBER OF SEQ ID NOS: 42  
; SOFTWARE: PatentIn version 3.0  
; SEQ ID NO 31  
; LENGTH: 936  
; TYPE: DNA

```
; ORGANISM: Archaeoglobus fulgidus
US-09-903-410-31

Query Match      22.0%; Score 28.4; DB 3; Length 936;
Best Local Similarity 56.4%; Pred. No. 8.4;
Matches 53; Conservative 0; Mismatches 41; Indels 0; Gaps 0;

Qy      2  GACCTTCTGCTCGTAGCGATTACTTCGAGCATTTACTGACGACAAAGACCCGACCGAGAT 61
        |||||  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
Db      718 GACCTTGAGAACTTACCTCTCGGCTGATCATACCGCGAATACGACCGCGTGAGAGAT 777

Qy      62  GGTTCGGGGTCTTTTGTGTGTGTGTGTGTGTGTGTGTGTGTGTGTGTGTGTGTGTGT 95
        |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
Db      778 GAAGGAGAGTTTTTCGGGCAGATGCTGAGAGAG 811
```

Search completed: March 7, 2006, 13:34:34  
Job time : 358.047 secs